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UTILITY PATENT APPLICATION TRANSMITTAL FORM (only for new nonprovisional applications under 37 CFR 1.53(b)

ASSISTANT COMMISSIONER FOR PATENTS Washington, D.C. 20231

BOX: PATENT APPLICATION

SIR:



Transmitted herewith for filing is the patent application (including Cover Pages and PCT Publication documents, Specification, Claims, and Abstract (84 pages) of:

Inventors: Dennis Gonsalves and Sheng-Zhi Pang

: TOSPOVIRUS RESISTANCE IN PLANTS For

**If a CONTINUING APPLICATION, please mark where appropriate and supply the requisite information below and in a preliminary amendment:

[X] continuation [] divisional [] Continuation-In-Part (CIP) of prior application Serial No. 08/495,484

Prior application information: Examiner: E. McElwain

Art Unit : 1649

Enci	osed are:							
[X]	5 sheets of informal drawings.							
[]	Signed Combined Declaration and Power of Attorney (pages).							
[X]	Copy of signed Combined Declaration and Power of Attorney (2 pages) from a prior application (1.63(d) (for continuation/divisional).							
[]	Signed statement deleting inventor(s) named in prior application (pages) (1.63(d)(2) and 1.33(b)).							
[X]	Incorporation By Reference : The entire disclosure of the prior application, from which a copy of the oath or declaration is supplied herewith, is considered as being part of the disclosure of the enclosed application and is hereby incorporated by reference therein.							
[]	Assignment (pages) of the invention to							
[]	Assignment Transmittal Letter.							
[]	Certified copy of a foreign priority document.							
[]	Associate power of attorney.							
[]	Verified statement to establish small entity status (pages) (newly signed or copy filed in prior application).							

- [X] Preliminary Amendment (11 pages).
- [X] Information Disclosure Statement, form PTO-1449 (5 pages) (in duplicate) and 39 references (not attached).
- [X] Sequence Listing (21 pages).
- [X] Statement in Accordance with 37 CFR §§ 1.821(f) and 1.825(b) and computer readable 3.5" Diskette.
- [X] A self-addressed, prepaid postcard acknowledging receipt.

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TOTAL CLAIMS	15 - 20 =	0				
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MULTIPLE DEPENDENT CLAIM PRESENTED						

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M2	ΔΤ	T	FNTITV

SMALL ENTITY						
RATE	FEE					
XXXX	\$380					
x 9=	\$					
x 39=	\$					
x130 =	\$					
TOTAL	\$					

LARGE ENTITY

<u>OR</u>	RATE	FEE
<u>OR</u>	XXXX	\$760
<u>OR</u>	x 18 =	\$
<u>OR</u>	X78 =	\$156
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- [] Please charge my Deposit Account No. ____ in the amount of \$____ . A duplicate copy of this sheet is enclosed.
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- [X] Address all future communications to:

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Date: October 22, 1999

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Applicants	:	Gonsalves et al.) Examiner:) To Be Assigned
Serial No.	:	Continuation of 08/495,484) Art Unit:
Filed	:	Herewith) To Be Assigned
For	:	TOSPOVIRUS RESISTANCE IN PLANTS))

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents Washington D.C. 20231

Sir:

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CHENT CONTROL OF THE REAL

Please amend the above-identified application as follows:

In the Title:

Please change the title to read -- TOSPOVIRUS RESISTANCE IN PLANTS--.

In the Specification:

On page 1, immediately below the Title of the Invention, please insert -- This application is a continuation of U.S. Patent Application Serial No. 08/495,484 filed September 25, 1995, as a national stage application of PCT Application No. PCT/US94/01046 filed January 27, 1994, which claims the priority benefit of U.S. Patent Application Serial No. 08/010,410 filed January 29, 1993.--

On page 3, line 18, change "in vivo" to --in vivo--.

On page 4, line 25 "change "viron" to --virion-- (each occurrence).

On page 4, line 27, change "viron" to --virion--.

On page 5, line 13, change "MK₂HPO₄" to --M K₂HPO₄--.

On page 5, lines 19-20, change "weight. The" to --weight, the--.

On page 6, line 23, before "complementary", insert -- (SEQ. ID. No. 1) --.

On page 6, line 24, before "of", insert -- (SEQ. ID. No. 2) --.

On page 6, line 27, change "clones" to --clone--.

On page 6, line 32, before "(also", insert -- (SEQ. ID. No. 3) --.

On page 7, line 9, before "(also", insert -- (SEQ. ID. No. 4) --.

On page 9, line 7, after the word "sequence" but before the ":", please insert -- (SEQ. ID. No. 5)--.

Please delete lines 8-35 on page 9 and lines 1-17 on page 10, and insert the following:

-- CAAGTTGAAA GCAACAACAG AACTGTAAAT TCTCTTGCAG TGAAATCTCT GCTCATGTCA 60 GCAGAAAACA ACATCATGCC TAACTCTCAA GCTTCCACTG ATTCTCATTT CAAGCTGAGC 120 CTCTGGCTAA GGGTTCCAAA GGTTTTGAAG CAGGTTTCCA TTCAGAAATT GTTCAAGGTT 180 GCAGGAGATG AAACAAACAA AACATTTTAT TTATCTATTG CCTGCATTCC AAACCATAAC 240 AGTGTTGAGA CAGCTTTAAA CATTACTGTT ATTTGCAAGC ATCAGCTCCC AATTCGCAAA 300 TGCAAAGCTC CTTTTGAATT ATCAATGATG TTTTCTGATT TAAAGGAGCC TTACAACATT 360 GTTCATGACC CTTCATACCC CAAAGGATCG GTTCCAATGC TCTGGCTCGA AACTCACACA 420 TCTTTGCACA AGTTCTTTGC AACTAACTTG CAAGAAGATG TAATCATCTA CACTTTGAAC 480 AACCTTGAGC TAACTCCTGG AAAGTTAGAT TTAGGTGAAA GAACCTTGAA TTACAGTGAA 540 GATGCCTACA AAAGGAAATA TTTCCTTTCA AAAACACTTG AATGTCTTCC ATCTAACACA 600 CAAACTATGT CTTACTTAGA CAGCATCCAA ATCCCTTCAT GGAAGATAGA CTTTGCCAGA 660 GGAGAAATTA AAATTTCTCC ACAATCTATT TCAGTTGCAA AATCTTTGTT AAAGCTTGAT 720 TTAAGCGGGA TCAAAAAGAA AGAATCTAAG GTTAAGGAAG CGTATGCTTC AGGATCAAAA 780 840 AATTATTTCT CTGTTTGTCA TCTCTTTCAA ATTCCTCCTG TCTAGTAGAA ACCATAAAAA 900 CAAAAAATAA AAATGAAAAT AAAATTAAAA TAAAATAAAA TCAAAAAAATG AAATAAAAAC 960 AACAAAAAT TAAAAAACGA AAAACCAAAA AGACCCGAAA GGGACCAATT TGGCCAAATT 1020 1080 TTTATTTTA TTTTATTTTT ATTTTATTTA TTTTTTGTTT TCGTTGTTTT TGTTATTTTA 1140 TTATTTATTA AGCACACAC ACAGAAAGCA AACTTTAATT AAACACACTT ATTTAAAATT 1200 TAACACTA AGCAAGCACA AGCAATAAAG ATAAAGAAAG CTTTATATAT TTATAGGCTT 1260 TTTTATAATT TAACTTACAG CTGCTTTCAA GCAAGTTCTG CGAGTTTTGC CTGCTTTTTA 1320 ACCCGAACA TTTCATAGAA CTTGTTAAGA GTTTCACTGT AATGTTCCAT AGCAACACTC 1380 CCTTTAGCAT TAGGATTGCT GGAGCTAAGT ATAGCAGCAT ACTCTTTCCC CTTCTTCACC 1440 TGATCTTCAT TCATTTCAAA TGCTTTGCTT TTCAGCACAG TGCAAACTTT TCCTAAGGCT 1500 TCCTTGGTGT CATACTTCTT TGGGTCGATC CCGAGGTCCT TGTATTTTGC ATCCTGATAT 1560 ATAGCCAAGA CAACACTGAT CATCTCAAAG CTATCAACTG AAGCAATAAG AGGTAAGCTA 1620 CCTCCCAGCA TTATGGCAAG TCTCACAGAC TTTGCATCAT CGAGAGGTAA TCCATAGGCT 1680 TGAATCAAAG GATGGGAAGC AATCTTAGAT TTGATAGTAT TGAGATTCTC AGAATTCCCA 1740 GTTTCTTCAA CAAGCCTGAC CCTGATCAAG CTATCAAGCC TTCTGAAGGT CATGTCAGTG 1800 CCTCCAATCC TGTCTGAAGT TTTCTTTATG GTAATTTTAC CAAAAGTAAA ATCGCTTTGC 1860 TTAATAACCT TCATTATGCT CTGACGATTC TTTAGGAATG TCAGACATGA AATAACGCTC 1920 ATCTTCTTGA TCTGGTCGAT GTTTTCCAGA CAAAAAGTCT TGAAGTTGAA TGCTACCAGA 1980 TTCTGATCTT CCTCAAACTC AAGGTCTTTG CCTTGTGTCA ACAAAGCAAC AATGCTTTCC 2040 TTAGTGAGCT TAACCTTAGA CATGATGATC GTAAAAGTTG TTATATGCTT TGACCGTATG 2100 TAACTCAAGG TGCGAAAGTG CAACTCTGTA TCCCGCAGTC GTTTCTTAGG TTCTTAATGT 2160 GATGATTTGT AAGACTGAGT GTTAAGGTAT GAACACAAAA TTGACACGAT TGCTCT 2216 --

On page 10, line 21, after "783" but before the ":", please insert --(SEQ. ID. No. 7)--.

Please delete lines 22-39 on page 10 and lines 1-18 on page 11, and insert the following:

-- Gln Val Glu Ser Asn Asn Arg Thr Val Asn Ser Leu Ala Val Lys Ser Leu Leu Met Ser Ala Glu Asn Asn Ile Met Pro Asn Ser Gln Ala Ser 25

Thr Asp Ser His Phe Lys Leu Ser Leu Trp Leu Arg Val Pro Lys Val Asn Ser Glu Leu Lys Gln Val Ser Ile Gln Lys Leu Phe Lys Val Asn Gly Asp Glu Thr Asn Lys Thr Phe Tyr Leu Ser Ile Ala Cys Ile Pro Asn His Asn 65

Ser Val Glu Thr Ala Leu Asn Ile Thr Val Ile Cys Lys His Gln Leu As Gly Asp Glu Rer Val Glu Thr Ala Leu Asn Ile Thr Val Ile Cys Lys His Gln Leu As Glu As Glu

Pro Ile Arg Lys Cys Lys Ala Pro Phe Glu Leu Ser Met Met Phe Ser Asp Leu Lys Glu Pro Tyr Asn Ile Val His Asp Pro Ser Tyr Pro Lys Gly Ser Val Pro Met Leu Trp Leu Glu Thr His Thr Ser Leu His Lys 130 135 Phe Phe Ala Thr Asn Leu Gln Glu Asp Val Ile Ile Tyr Thr Leu Asn 145 Asn Leu Glu Leu Thr Pro Gly Lys Leu Asp Leu Gly Glu Arg Thr Leu 170 165 Asn Tyr Ser Glu Asp Ala Tyr Lys Arg Lys Tyr Phe Leu Ser Lys Thr 185 Leu Glu Cys Leu Pro Ser Asn Thr Gln Thr Met Ser Tyr Leu Asp Ser 195 200 Ile Gln Ile Pro Ser Trp Lys Ile Asp Phe Ala Arg Gly Glu Ile Lys 215 Ile Ser Pro Gln Ser Ile Ser Val Ala Lys Ser Leu Leu Lys Leu Asp 225 230 Leu Ser Gly Ile Lys Lys Glu Ser Lys Val Lys Glu Ala Tyr Ala 250 Ser Gly Ser Lys

On page 11, line 21, after the word "gene" (second occurrence) but before the ".", please insert --(SEQ. ID. No. 6)--.

On page 12, line 20, after the word "gene" but before the ".", please insert -- (SEQ. ID. No. 8)--.

On page 13, line 9, after the word "is" and before ":", please insert --(SEQ. ID. No. 9)--.

On page 16, line 23, change "in vitro" to --in vitro--.

On page 16, line 33, change "a nd" to --and--.

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On page 18, line 31, change "teh" to --the--.

On page 19, line 26, change "TSWV-BI" to --TSWV-BL--.

On page 21, line 32, change "viron" to --virion--.

On page 22, line 7, change "viron" to --virion--.

On page 22, line 28, change "symptomless" to --Symptomless--.

On page 22, line 33, change "cold" to --could--.

On page 23, line 18, change "viron" to --virion--.

On page 24, line 8, change "symptom less" to --symptomless--.

On page 26, line 22, before the ",", insert -- (SEQ. ID. No. 10) --.

On page 26, line 33, before "complementary", insert -- (SEQ. ID. No. 11) --.

On page 27, line 1, change "anshor" to --anchor--.

On page 27, line 2, before "that", insert -- (SEQ. ID. No. 17) --.

On page 27, line 4, change "amplified fragement" to --amplified fragment--.

On page 27, line 7, before "close", insert -- (SEQ. ID. No. 16) --.

On page 27, line 28, after the word "is" but before the ":", please insert -- (SEQ. ID. No. 14)--.

On page 29, line 22, after the word "are" but before the ":", please insert -- (SEQ. ID. No. 12)--.

Please delete lines 23-38 on page 29, lines 1-44 on page 30, and lines 1-3 on page 31, and insert the following:

-- Met Ser Ser Gly Val Tyr Glu Ser Ile Ile Gln Thr Lys Ala Ser Val

Trp Gly Ser Thr Ala Ser Gly Lys Ser Ile Val Asp Ser Tyr Trp Ile
20 25 30

Tyr Glu Phe Pro Thr Gly Ser Pro Leu Val Gln Thr Gln Leu Tyr Ser 35 40 45

Asp Ser Arg Ser Lys Ser Ser Phe Gly Tyr Thr Ser Lys Ile Gly Asp 50 55 60

Ile Pro Ala Val Glu Glu Glu Ile Leu Ser Gln Asn Val His Ile Pro 65 70 75 80

Val Phe Asp Asp Ile Asp Phe Ser Ile Asn Ile Asn Asp Ser Phe Leu 85 90 95

Ala Ile Ser Val Cys Ser Asn Thr Val Asn Thr Asn Gly Val Lys His
100 105 110

Gln Gly His Leu Lys Val Leu Ser Leu Ala Gln Leu His Pro Phe Glu 115 120 125

Pro Val Met Ser Arg Ser Glu Ile Ala Ser Arg Phe Arg Leu Gln Glu 130 135 140

Ser Leu Ser Cys Val Lys Glu His Thr Tyr Lys Val Glu Met Ser His

Asn Gln Ala Leu Gly Lys Val Asn Val Leu Ser Pro Asn Arg Asn Val 185 His Glu Trp Leu Tyr Ser Phe Lys Pro Asn Phe Asn Gln Ile Glu Ser 200 Asn Asn Arg Thr Val Asn Ser Leu Ala Val Lys Ser Leu Leu Met Ala 215 Thr Glu Asn Asn Ile Met Pro Asn Ser Gln Ala Phe Val Lys Ala Ser Thr Asp Ser His Phe Lys Leu Ser Leu Trp Leu Arg Ile Pro Lys Val 250 245 Leu Lys Gln Ile Ala Ile Gln Lys Leu Phe Lys Phe Ala Gly Asp Glu Thr Gly Lys Ser Phe Tyr Leu Ser Ile Ala Cys Ile Pro Asn His Asn 275 Ser Val Glu Thr Ala Leu Asn Val Thr Val Ile Cys Arg His Gln Leu 295 Pro Ile Pro Lys Ser Lys Ala Pro Phe Glu Leu Ser Met Ile Phe Ser 310 315 Asp Leu Lys Glu Pro Tyr Asn Thr Val His Asp Pro Ser Tyr Pro Gln Arg Ile Val His Ala Leu Leu Glu Thr His Thr Ser Phe Ala Gln Val Leu Cys Asn Lys Leu Gln Glu Asp Val Ile Ile Tyr Thr Ile Asn Ser Pro Glu Leu Thr Pro Ala Lys Leu Asp Leu Gly Glu Arg Thr Leu Asn Tyr Ser Glu Asp Ala Ser Lys Lys Lys Tyr Phe Leu Ser Lys Thr Leu Glu Cys Leu Pro Val Asn Val Gln Thr Met Ser Tyr Leu Asp Ser Ile Gln Ile Pro Ser Trp Lys Ile Asp Phe Ala Arg Gly Glu Ile Arg Ile Ser Pro Gln Ser Thr Pro Ile Ala Arg Ser Leu Leu Lys Leu Asp Leu Ser Lys Ile Lys Glu Lys Lys Ser Leu Thr Trp Glu Thr Ser Ser Tyr 455 Asp Leu Glu 465

On page 31, line 4, after the word "and", please insert -- (SEQ. ID. No. 13)--.

On page 31, line 41, after the word "is" but before the ":", please insert -- (SEQ. ID. No. 15)--.

Please delete lines 42-43 on page 31 and lines 1-16 on page 32, and replace with the following:

-- TTATGCAACA CCAGCAATCT TGGCCTCTTT CTTAACTCCA AACATTTCAT AGAATTTGTC 60 AAGATTATCA CTGTAATAGT CCATAGCAAT GCTTCCCTTA GCATTGGGAT TGCAAGAACT 120 AAGTATCTTG GCATATTCTT TCCCTTTGTT TATCTGTGCA TCATCCATTG TAAATCCTTT 180 GCTTTTAAGC ACTGTGCAAA CCTTCCCCAG AGCTTCCTTA GTGTTGTACT TAGTTGGTTC 240 AATCCCTAAC TCCTTGTACT TTGCATCTTG ATATATGGCA AGAACAACAC TGATCATCTC 300 GAAGCTGTCA ACAGAAGCAA TGAGAGGGAT ACTACCTCCA AGCATTATAG CAAGTCTCAC 360 AGATTTTGCA TCTGCCAGAG GCAGCCCGTA AGCTTGGACC AAAGGGTGGG AGGCAATTTT 420 TGCTTTGATA ATAGCAAGAT TCTCATTGTT TGCAGTCTCT TCTATGAGCT TCACTCTTAT 480 CATGCTATCA AGCCTCCTGA AAGTCATATC CTTAGCTCCA ACTCTTTCAG AATTTTTCTT 540 TATCGTGACC TTACCAAAAG TAAAATCACT TTGGTTCACA ACTTTCATAA TGCCTTGGCG 600 ATTCTTCAAG AAAGTCAAAC ATGAAGTGAT ACTCATTTTC TTAATCAGGT CAAGATTTTC 660 CTGACAGAAA GTCTTAAAGT TGAATGCGAC CTGGTTCTGG TCTTCTTCAA ACTCAACATC 720 TGCAGATTGA GTTAAAAGAG AGACAATGTT TTCTTTTGTG AGCTTGACCT TAGACAT 777

The complementary nucleic acid molecule has a nucleotide sequence as shown in SEQ. ID. No. 19.--.

On page 32, line 24, change "vial" to --viral--.

On page 33, line 29, change "the" to -- The--.

On page 33, line 32, change "differs" to --differ--.

On page 34, line 17, change "exactly same positions" to --exact same position--.

On page 35, line 12, change "event." to --event,--.

On page 35, line 13, change "Tospovirus" to --Tospovirus--.

On page 36, line 10, change "generated" to --generate--.

On page 36, line 27, change "Np" to --NP--.

On page 39, line 16, after "virus" insert --was--.

On page 39, line 17, change ", and is" to -- and was--.

On page 40, line 25, change "resistant" to --resistance--.

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On page 41, line 3, before "and", insert -- (SEQ. ID. No. 20) --.
               On page 41, line 4, before ",", insert -- (SEQ. ID. No. 21) --.
               On page 41, line 10, after "plants" insert --that--.
               On page 41, line 11, change "greenhouse. All" to --greenhouse, all--.
               On page 43, line 9, after "appearance" insert --.--
               On page 43, line 15, change "pants" to --plants--.
               On page 45, line 19, change "vecor" to -- vector --.
               On page 45, line 33, before ",", insert -- (SEQ. ID. No. 22) --.
               On page 45, line 35, before ",", insert -- (SEQ. ID. No. 23) --.
               On page 46, line 31, before "was", insert -- (SEO, ID, No. 24) --.
               On page 47, line 11, change "Hewlet Scanjet" to --Hewlett Scanjet<sup>TM</sup>---.
               On page 51, line 2, change "Tospoviruses" to -- Tospoviruses--.
               On page 53, line 11, change "Tospovirus" to --Tospovirus--.
               On page 53, line 14, change "realated" to --related--.
               On page 53, line 16, change "RNE" to -- RNA --.
               On page 53, line 28, change "generated" to --generated--.
               On page 54, line 2, before ",", insert -- (SEQ. ID. No. 25) --.
               On page 54, line 4, before "for", insert -- (SEQ. ID. No. 26) --.
               On page 54, line 6, before "for", insert -- (SEQ. ID. No. 27) --.
               On page 54, line 10, before "which", insert -- (SEO, ID, No. 28) --.
               On page 54, line 12, before "for", insert -- (SEQ. ID. No. 29) --.
               On page 54, line 15, before "for", insert -- (SEQ. ID. No. 30) -- and change
"untranslatablesecond" to -- untranslatable second --.
               On page 54, line 33, change "expressin" to --expression--.
               On page 55, line 23, change "fragemnts" to --fragments--.
               On page 55, line 32, change "fragements" to --fragments--.
               On page 55, line 30, change "prodeing" to -- producing --.
               On page 55, line 34, change "expressin" to -- expression --.
               On page 56, line 4, change "trnscription" to --transcription--.
               On page 56, line 6, change "pant" to -- plant -- and change "resluting" to --
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On page 56, line 32, change "variojs" to -- various --.

On page 57, line 23, change "portin" to -- portion --.

resulting --.

Please delete the sequence listing from the specification, page 57, line 26 to page 75, line 9 and insert the new sequence listing (attached hereto) as a separately paginated document which is part of the present application.

In the Claims:

Please delete claims 1-3 and add new claims 4-18 as follows:

- --4. An isolated DNA molecule capable of transcription to a messenger RNA which is modified from a form encoding a nucleocapsid protein of an L serogroup *Tospovirus* so that it does not translate to the nucleocapsid protein, wherein, when the DNA molecule is transformed into a plant cell, it is capable of being transcribed into messenger RNA which exists at low level density readings of 15-50 as measured using a Hewlett ScanJet and Image Analysis Program.--
- --5. A DNA molecule according to claim 4, wherein the L serogroup *Tospovirus* is selected from the group consisting of TSWV-10W and TSWV-BL.--
- --6. A recombinant DNA expression system comprising an expression vector into which is inserted a heterologous DNA molecule according to claim 4.--
- --7. A plant cell transformed with a heterologous DNA molecule according to claim 4.--
- --8. A transgenic plant containing the DNA molecule according to claim 4.--
- --9. A method of treating a plant cell comprising:
 transforming a plant cell with a DNA molecule according to claim 4
 and

transcribing the DNA molecule under conditions effective to maintain the messenger RNA in the plant cell at low level density readings of 15-50, as measured

using the Hewlett ScanJet and Image Analysis Program, wherein the plant cell acquires resistance to an L serogroup *Tospovirus*.--

--10. A method of imparting to a plant cell resistance to infection by an I serogroup *Tospovirus*, said method comprising:

transforming a plant cell with a DNA molecule encoding a nucleocapsid protein of an L serogroup *Tospovirus*, wherein the DNA molecule is expressed to produce an ELISA level of OD405nm = 0.50 to 1.00 of the nucleocapsid protein in the plant cell, as measured using an antibody raised against the nucleocapsid protein of *Tospovirus* isolate TSWV-BL, and the plant cell acquires resistance to an I serogroup *Tospovirus*.--

- --11. A method according to claim 10, wherein the L serogroup *Tospovirus* is selected from the group consisting of TSWV-10W and TSWV-BL.--
- --12. A method according to claim 10, wherein the I serogroup *Tospovirus* is selected from the group consisting of INSV-Beg and INSV-LI.--
- --13. A method of imparting to a plant cell resistance to infection by L serogroup *Tospoviruses*, said method comprising:

transforming a plant cell with a DNA molecule encoding a nucleocapsid protein of an L serogroup *Tospovirus*, wherein the DNA molecule is expressed to produce an ELISA level of OD 405nm = 0.02 to 0.20 of the nucleocapsid protein in the plant cell, as measured using an antibody raised against the nucleocapsid protein of *Tospovirus* isolate TSWV-BL, and the plant cell acquires resistance to the L serogroup *Tospovirus* and a second L serogroup *Tospovirus*.--

- --14. A method according to claim 13, wherein the L serogroup *Tospovirus* is selected from the group consisting of TSWV-10W and TSWV-BL.--
- --15. A method of imparting to a plant cell resistance to infection to a *Tospovirus* comprising:

transforming a plant cell with a DNA molecule encoding a nucleocapsid protein or polypeptide of a serogroup L *Tospovirus* under conditions effective to render the plant cell resistant to infection by serogroup L *Tospovirus* isolates and a serogroup 2 *Tospovirus* isolate.--

- --16. A method according to claim 15, wherein the serogroup 2 *Tospovirus* isolate is a TSWV-B *Tospovirus* isolate.--
- --17. A transgenic plant containing a heterologous DNA molecule encoding a nucleocapsid protein or polypeptide of a serogroup L *Tospovirus*, wherein the transgenic plant, upon challenge with both a *Tospovirus* isolate belonging to serogroup L and a *Tospovirus* isolate belonging to serogroup 2, exhibits resistance to both the L serogroup *Tospovirus* isolate and the serogroup 2 *Tospovirus* isolate.--
- --18. A transgenic plant according to claim 17, wherein the serogroup 2 *Tospovirus* isolate is a TSWV-B *Tospovirus* isolate.--

Date: October 22, 1999

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TOMATO SPOTTED WILT VIRUS

Viruses in the *Tospovirus* genus infect a wide variety of plant species, particularly tobacco, peanut, vegetables and ornamental plants. Two virus species, tomato spotted wilt virus (TSWV) and impatiens necrotic spot virus (INSV) are recognized within the Tospovirus genus.

Tomato Spotted Wilt Virus (TSWV) is unique among plant viruses in that the nucleic acid-protein complex is covered by a lipoprotein envelope and it is the only thrip transmitted virus. This virus has recently been classified as the Tospovirus genus of the *Bunyaviridae* family. TSWV virions contain a 29K nucleocapsid protein ("NP" or "N"), two membrane-associated glycoproteins (58K and 78K) and a large 200K protein presumably for the viral transcriptase [see J. Gen. Virol. 71:2207 (1991); Virol. 56:12 (1973); and J. Gen. Virol. 36:267 (1977)].

15 The virus genome consists of three negative-strand (-) RNAs designated L RNA (8900 nucleotides), M RNA (5400 nucleotides) and S RNA (2900 nucleotides) [see J. Gen. Virol. 36:81 (1977); J. Gen. Virol. 53:12 (1981); and J. Gen. Virol. 70:3469 (1989)], each of which is encapsulated by the NP. The partial or full-length sequences of S RNAs from three TSWV isolates reveals the presence of two open reading frames (ORF) with an

isolates reveals the presence of two open reading frames (ORF) with an ambisense gene arrangement [see J. Gen Virol. 71:1 (1990) and J. Gen. Virol. 72:461 (1991)]. The larger open reading frame is located on the viral RNA strand and has the capacity to encode a 52K nonstructural protein. The smaller ORF is located on the viral complementary RNA strand and is translated through a subgenomic RNA into the 29K NP.

The ambisense coding strategy is also characteristic of the TSWV M RNA, with the open reading frames encoding the 58K and 78K membrane-associated glycoproteins. The TSWV L RNA has been sequenced to encode a large 200K protein presumably for the viral transcriptase.

Two TSWV serogroups, "L" and "l", have been identified and characterized based on serological analysis of the structural proteins and morphology of cytopathic structures [see J. Gen Virol. 71:933 (1990) and Phytopathology 81:525 (1991)]. They have serologically conserved G1 and G2 glycoproteins, but the NP of the "I" serogroup is

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serologically distinct from that of the "L" serogroup. Comparison of the NP between the "L" and "I" serogroups has shown 62% and 67% identities at nucleotide and amino acid levels, respectively [see J. Gen. Virol. 72:2597 (1991)].

TSWV has a wide host range, infecting more than 360 plant species of 50 families and causes significant economic losses to vegetables and ornamental plants worldwide. The "L" serogroup has been found extensively in field crops such as vegetables and weeds, while the "I" serogroup has been largely confined to ornamental crops.

A cucurbit isolate has recently been identified [see Plant Disease 68:1006 (1984)] as a distinct isolate because it systemically infects watermelon and other curcurbits and its NP is serologically unrelated to that of either serogroup. Although the spread of the TSWV disease can sometimes be reduced by breeding resistant plants or using non-genetic approaches, complete control of the disease by these conventional methods has generally proven to be difficult [see Plant Disease 73:375 (1989)].

Since 1986, numerous reports have shown that transgenic plants with the coat protein (CP) gene of a virus are often resistant to infection by that virus. This phenomenon is commonly referred to as coat protein-mediated protection (CPMP). The degree of protection ranges from delay in symptom expression to the absence of disease symptoms and virus accumulation. Two recent independent reports [see Biol. Technology 9:1363(1991) and Mol. Plant-Microbe Interact.

5:34 (1992)] showed that transgenic tobacco plants expressing the nucleocapsid protein (NP) gene of TSWV are resistant to infection by the homologous isolate. However, since TSWV is widespread with many biologically diverse isolates, it is very important to test the effectiveness of the transgenic plants to resist infections by different TSWV isolates. The findings of the present invention expand on those of the previous reports by demonstrating that transgenic plants according to the present invention showed resistance to two heterologous isolates of the "L" serogroup and an isolate of the "I" serogroup. We also show that resistance to the two heterologous isolates of the "L" serogroup was mainly found in plants accumulating very low, if any,

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levels of NP, while transgenic plants that accumulated high levels of NP were resistant to the isolate of the "I" serogroup.

However, no resistance was observed to a Brazilian isolate, although the plants that accumulated high levels of the N protein did display a delay in symptom expression. This Brazilian isolate, designated TSWV-B has the N protein that was serologically distinct from the "L" and "I" serogroups and biologically differs from a curcurbit isolate in that the TSWV-B does not systemically infect melons or squash. Therefore, one aspect of the present invention is to characterize the TSWV-B by cloning and sequencing of its S RNA and comparisons with the published sequences of other TSWV isolates.

Various aspects of the present invention will become readily apparent from the detailed description of the present invention including the following example, figures and data.

15 In the Figures;

Fig. 1 depicts the strategy for cloning the NP gene from viral RNA according to the present invention;

Fig. 2 depicts the In vivo transient expression of the nucleocapsid protein (NP) gene of tomato spotted wilt virus according to the present invention in tobacco protoplasts;

Fig. 3 depicts the location of the sequenced cDNA clones in the TSWV-B S RNA according to the present invention;

Fig. 4 depicts a dendogram showing relationships among TSWV isolates according to the present invention;

Fig. 5 depicts the serological relationship of TSWV isolates described herein;

Fig. 6 depicts the correlation of the level of nucleocapsid protein (NP) accumulation in transgenic plants with the degree of resistance to TSWV isolates;

Fig. 7 depicts the TSWV-BL N coding sequences introduced into transgenic plants in accordance with one aspect of the present invention; and

Fig. 8 depicts the TSWV-BL half N gene fragments introduced into plants in accordance with one aspect of the present invention.

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More specifically, figure 2 depicts transient expression of the NP gene in which the constructs were transferred into tobacco mesophyll protoplasts using polyethylene glycol (PEG). The transformed protoplasts were subsequently incubated for two days for the expression of the NP gene. Proteins were extracted from the protoplasts and tested for the NP by double antibody sandwich enzymelinked immunosorbent assay (DAS-ELISA) using antibodies against the TSWV NP. NP and NP+ represent the protoplasts transformed with plasmids pBI525-NP- and pBI525-NP+, respectively. Concentration of the antibodies for coating: 5 μg/ml: dilution of the enzyme conjugate: 1:250. Data were taken 30, 60 and 90 min. after addition of substrate.

In figure 3, the five overlapping cDNA clones are shown to scale below a S RNA map of TSWV-B. These clones were synthesized with random primers from double-stranded RNA isolated from N. benthamiana plants infected with TSWV-B.

In figure 4, the sequences were compared using the pileup program of the GCG Sequence analysis software package. Horizontal lines are proportional to the genetic distance while vertical lines are of arbitrary length and have no significance.

More specifically, in figure 5, N. benthamiana Domin. were infected with TSWV isolates [TSWV-BL (a lettuce isolate), Arkansas, 10W pakchoy (TSWV-10W), Begonia, and Brazil (TSWV-B)). An infected leaf disc (0.05 gram) was ground in 12 ml of the enzyme conjugate buffer and analyzed by DAS-ELISA using antibodies raised against

- TSWV-BL viron (BL viron), or the NP of TSWV-BL (BL-NP), or TSWV-I (I-NP). Concentration of antibodies for coating were 1μg/ml; dilution of conjugates were 1:2000 for BL viron, 1:250 for BL-NP, and 1:1000 for I-NP. The results were taken after 10 minutes (BL), 50 minutes (BL-NP), or 30 minutes after adding substrate.
- With regard to figure 6, transgenic plants were assayed in DAS-ELISA for NP accumulation with antibodies raised against the NP of TSWV-BL. Plants were read 150 min. after adding substrate and the transgenic plants were grouped into four categories: OD_{405nm} smaller than 0.050, OD_{405nm} between 0.050 to 0.200, OD_{405nm} between 0.200

to 0.400, and OD405nm greater than 0.400. The OD405nm readings of

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control NP (-) plants were from zero to 0.05. The same plants were challenged with either the Arkansas (Ark) and 10W pakchoy (10W) isolates or the Begorila isolate and the susceptibility of each plant was recorded about 12 days after inoculation. The results were pooled from filty-one R₁ NP (+) plants inoculated with the Arkansas and 10W pakchoy isolates and one hundred thirty-nine R₁ NP(+) plants inoculated with the Begorila isolate. Numbers above bars represent total numbers of R₁ NP(+) plants tested.

EXAMPLE I

10 Isolation of TSWV-BL RNAs:

The TSWV-BL isolate was purified from Datura stramonium L. as follows: the infected tissues were ground in a Waring Blender for 45 sec with three volumes of a buffer (0.033 M KH2PO4, 0.067 MK2HPO4, 0.01 M Na2SO3). The homogenate was filtered through 4 layers of cheesecloth moistened with the above buffer and centrifuged at 7,000 rpm for 15 min. The pellet was resuspended in an amount of 0.01 M Na2SO3 equal to the original weight of tissue and centrifuged again at 8,000 rpm for 15 min. After the supernatant was resuspended in an amount of 0.01 M Na2SO3 equal to 1/10 of the original tissue weight.

The virus extract was centrifuged at 9,000 rpm for 15 min. and the supernatant was carefully loaded on a 10-40% sucrose step gradient made up in 0.01 M Na₂SO₃. After centrifugation at 23,000 rpm for 35 min., the virus zone (about 3 cm below meniscus) was collected and diluted with two volumes of 0.01 M Na₂SO₃. The semi-purified virus was pelleted at 27,000 rpm for 55 min.

EXAMPLE II

Purification of TSWV and viral RNAs:

The TSWV-BL isolate [see Plant Disease 74:154 (1990)] was purified from *Datura stramonium* L, as described in Example I. The purified virus was resuspended in a solution of 0.04% of bentonite, 10 µg/ml of proteinase K, 0.1 M ammonium carbonate, 0.1% (w/v) of sodium diethyldithiocarbanate, 1 mM EDTA, and 1% (w/v) of sodium dodecyl sulfate (SDS), incubated at 65°C for 5 min., and immediately extracted from H₂O-saturated phenol, followed by another extraction

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with chloroform/isoamyl alcohol (24:1). Viral RNAs were precipitated in 2.5 volumes of ethanol and dissolved in distilled H₂O.

EXAMPLE III

cDNA and PCR-based NP gene cloning:

The first strand cDNA was synthesized from purified TSWV-BL RNAs using random primers as described by Gubler and Hoffman [see Gene 25:263 (1983)). The second strand was produced by treatment of the sample with RNase H/DNA polymerase. The resulting double-, stranded cDNA sample was size-fractionated by sucrose gradient centrifugation, methylated by EcoRI methylase, and EcoRI linkers were added. After digestion with EcoRI, the cDNA sample was ligated into the EcoRI site of pUC18, whose 5'-terminal phosphate groups were removed by treatment with calf intestinal alkaline phosphotase. E. coli DH5 α competent cells (Bethesda Research Laboratories) were transformed and clones containing TSWV cDNA inserts were first selected by plating on agar plates containing 50 µg/ml of ampicillin, IPTG, and X-gal. Plasmid DNAs from selected clones were isolated using an alkaline lysis procedure [see BRL Focus 11:7 (1989)], and the insert sizes were determined by EcoRI restriction enzyme digestion followed by DNA transfer onto GeneScreen Plus nylon filters (DuPont). Plasmid clones that contained a TSWV-BL S RNA cDNA insert were identified as described below by hybridizing against a 32P-labelled oligomer (AGCAGGCAAAACTCGCAGAACTTGC) complementary to the nucleotide sequence (GCAAGTTCTGCGAGTTTTGCCTGCT) of the TSWV-CPNH1 S RNA [see J. Gen. Virol. 71:001 (1990)]. Several clones were

25 CPNH1 S RNA [see J. Gen. Virol. 71:001 (1990)]. Several clones were identified and analyzed on agarose gels to determine the insert sizes. The clones pTSWVS-23 was found to contain the largest cDNA insert, about 1.7 kb in length.

The full-length NP gene was obtained by the use of polymerase chain reaction (PCR). First-strand cDNA synthesis was carried out at 37°C for 30 min. In a 20 µl reaction mixture using oligomer primer JLS90-46 (5'-> 3') AGCTAACCATGGTTAAGCTCACTAAGGAAAGC (also used to synthesize the nucleocapsid gene of TSWV-10W) which is complementary to the S RNA in the 5' terminus of TSWV NP gene

3.5 (nucleotide positions 2751 to 2773 of the TSWV-CPNH1). The reaction

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mixture contained 1.5 µg of viral RNAs,1 µg of the oligomer primer, 0.2 mM of each dNTP, 1X PCR buffer (the GeneAmp kit, Perkin-Elmer-Cetus). 20U of RNAs in Ribonuclease inhibitor (Promega), 2.5 mM of MgCl2, and 25U of AMV reverse transcriptase (Promega Corporation). The reaction was terminated by heating at 95°C for 5 min. and cooled on ice. Then 10 ul of the cDNA/RNA hybrid was used to PCR-amplify the NP gene according to manufacturer's instructions (Perkin-Elmer-Cetus) using 1 ug each of oligomer primers JLS90-46 and JLS90-47 (5'->3'), AGCATTCCATGGTTAACACACACTAAGCAAGCAC (also used to synthesize the nucleotide gene of TSWV-10W), the latter oligomer being identical to the S RNA in the 3' noncoding region of the gene (nucleotide positions 1919 to 1938 of the TSWV-CPNH1). A typical PCR cycle was 1 min. at 92°C (denaturing), 1 min. at 50°C (annealing), and 2 min. at 72°C (polymerizing). The sample was directly loaded and separated on a 1.2% agarose gel. The separated NP gene fragment was extracted from the agarose gel, ethanol-precipitated and dissolved in 20 µl of distilled H₂O.

EXAMPLE IV

Construction of plant expression and transformation vectors.

The gel-isolated NP gene fragment from Example III was digested with the restriction enzyme Ncol in 50 µl of a reaction buffer [50 mM Tris-HCI (pH 8.0), 10 mM MgCl₂, 0.1 M NaCl] at 37°C for 3 hours, and directly cloned into Ncol-digested plant expression vector pB1525. The resulting plasmids were identified and designated as pB1525-NP+ in the sense orientation relative to the cauliflower mosaic virus (CaMV) 35S promoter, and as pB1525-NP in the reverse orientation. The ability of this expression cassette to produce the NP was determined by transient expression of the NP gene in Nicotiana tobacum protoplasts, as described by Pang et al [see Gene 112:229 (1992)]. The expression cassette containing the NP gene was then excised from pB1525-NP+ by a partial digestion with HindIII/EcoRI (since the NP gene contains internal HindIII and EcoRI sites), and ligated into the plant transformation vector pBIN19 (Clontech Laboratories, Inc.) that had been cut with the same enzymes. The resulting vector, pBIN19-NP+ and the control plasmid pBIN19 were transferred to A. tumefaciens strain

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LBA4404, using the procedure described by Holsters et al [see Mol. Gen. Genet. 163:181 (1978)].

Nucleotide sequence analyses of the inserts in clones pTSWV-23 and Pb1525-NP+were determined using the dideoxyribonucleotide method, T7 polymerase (U.S. Biochemicals, SequenaseTM), and the double-stranded sequencing procedure described by Siemieniak et al [see Analyt. Biochem. 192:441 (1991)]. Nucleotide sequences were determined from both DNA strands and this information was compared with the published sequences of TSWV isolates CPNH1 using computer programs available from the Genetics Computer Group (GCG, Madison, WI).

Transient expression of the NP gene in tobacco protoplasts were also prepared. Plasmid DNAs for clones pTSWVS-23 and pUC18cpphas TSWV-NP (containing the PCR-engineered NP gene insert) were isolated using the large scale alkaline method. The PCR-engineered NP gene insert was excised from clone pBIS25-NP+ by Ncol digestion to take advantage of the available flanking oligomer primers for sequencing. The expression cassette pUC18cpphas is similar to pUC18cpexp except that it utilizes the poly(A) addition signal derived from the Phaseolus vulgaris seed storage gene phaseolin. These plasmid DNAs were subjected to two CsCl-ethidium bromide gradient bandings, using a Beckman Ti 70.1 fixed angle rotor. DNA sequences were obtained using dideoxyribonucleotides and the double-stranded plasmid DNA sequencing procedure described above. Nucleotide sequence reactions were electrophoresed on one-meter long thermostated (55°C) sequencing gels and nucleotide sequence readings averaging about 750 bp were obtained. Nucleotide sequences were determined from both DNA strands of both cloned inserts to ensure accuracy. Nucleotide sequence information from the TSWV-BL S RNA Isolate was compared as discussed below, with TSWV isolates CPNH1 and L3 using computer programs (GCG, Madison, WI).

The nucleotide and deduced amino acid sequences of cloned cDNA and PCR-engineered insert of TSWV-BL S RNA and their comparison with the nucleotide sequence of TSWV-CPHN1 S RNA are shown below.

3.5 The nucleotide sequence of the TSWV-BL S RNA clones pTSWVS-23

(TSWV-23) and pBI525-NP+ (TSWV-PCR) were obtained using the double-stranded dideoxynucleotide sequencing procedure of Siemieniak, and their sequences are compared with the relevant regions of the nucleotide sequence of the TSWV-CPNH1 S RNA reported in GeneBank

Accession No. D00645. The nucleotide sequence of TSWV-CPNH1 S RNA has been reported by De Haan (1990) and is represented by the following sequence:

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	CAÁGITGAAA GCAACAACAG ÁACIGIAAAT TCICTIGCAG IGAAATCICT	, 50
	GCTCÁTGTCA GCAGAÁÁCA ÁCATCATGCC TAACTCTCÁA GCTTCCACTG	100
10	ATTCTCATTT CAAGCTGAGC CTCTGGCTAA GGGTTCCAAA GGTTTTGAAG	150
	CAGGITICCA TICAGAAATT GIICAAGGIT GCAGGAGATG AAACAAACAA	200
	AACATTITAT TIATCIATIG CCIGCATICC AAACCATAAC AGIGIIGAGA	250
	CAGCITTAAA CATTACIGIT ATTIGCAAGC ATCAGCICCC AATTCGCAAA	300
	TGCAAAGCIC CITTIGAATT ATCAATGATG TITTCIGATT TAAAGGAGCC	350
15	TIACAACATT GITCATGACC CITCATACCC CAAAGGATCG GITCCAATGC	400
	TCTGGCTCGA AACTCACACA TCTTTGCACA AGTTCTTTGC AACTAACTTG	450
	CAÁGAAGAIG TAATCATCIA CACITIGAAC AACCITGAGC TAACICCIGG	500
	AAAGITAGAT TTAGGIGAAA GAACCITGAA TIACAGIGAA GATGCCTACA	550
	AAAGGAAATA TITCCITICA AAAACACITG AATGICITCC ATCIAACACA	600
20	CAAACTATGT CITACTTAGA CAGCATCCAA ATCCCTTCAT GGAAGATAGA	650
	CTTIGCCAGA GGAGAAATTA AAATTICICC ACAATCTATI TCAGITGCAA	700
	AATCITIGIT AAAGCITGAT TTAAGCGGGA TCAAAAAGAA AGAATCIAAG	750
	GITAAGGAAG CGTATGCITC AGGATCAAAA TAATCITGCT TIGICCAGCT	800
	TTTTCTAATT ATGITATGIT TATTTTCITI CTTTACITAT AATTATTTCT	850
25	CIGITIGICA TCICITICAA ATTCCICCIG TCIAGIAGAA ACCATAAAAA	900
	CAAAAATAA AAATGAAAAT AAAATTAAAA TAAAATAAAA TCAAAAAATG	1000
	AAATAAAAC AACAAAAAT TAAAAAACGA AAAACCAAAA AGACCCGAAA	1050
	GGGACCAATT TGGCCAAATT TGGGTTTTGT TTTTGTTTTT TGTTTTTTGT	1100
•	TITTIATITI TITTIATITI TATTITATITI TITATITITA TITTATITIT	1150
30	ATTTATTIA TITTTIGITT TCGITGITTT TGTTATTITA TIATTLATTA	1200
	AGCACAACAC ACAGAAAGCA AACITTAATT AAACACACIT ATTTAAAATT	1250
	INACACACIA AGCAAGCACA AGCAATAAAG ATAAAGAAAG CITTATATAT	1300
	TIATAGGCIT TTITATAATT TAACITACAG CIGCITICAA GCAAGITCIG	1350
	CGAGITITGC CIGCITITTA ACCCCGAACA TITCATAGAA CITGITAAGA	1400
3.5	GTTTCACTGT AATGTTCCAT AGCAACACTC CCTTTAGCAT TAGGATTGCT	1450

	GGAGCTAAGT	ATAGCAGCAT	ACTOTTTOCC	CTTCTTCACC	TGATCTTCAT	1500
	TCATTTCAAA '	TGCTTTGCTT	TTCAGCACAG	TGCAAACTTT	TCCTAAGGCT	1550
	TCCTTGGTGT	CATACTICIT	TGGGTCGATC	COGAGGICCT	TGTATTTTGC	1600
	ATCCIGATAT	ATAGCCÀÁGA	CAACACTGAT	CATCTCAAAG	CTATCAACIG	1650
5	AAGCAATAAG	AGGIAAGCIA	CCTCCCAGCA	TTATGGCAAG	TCTCACAGAC	1700
	TTTGCATCAT	OGÁGAGGÍTAÁ	TCCATAGGCT	TGAATCAAAG	GATGGGAAGC	1750
	AATCTTÁGAT	TIGATAGIÁT	TGAGATTCTC	AGAATTCCCA	GITICTICAA	1800
	CAAGCCTGAC	CCTGATCÁAG	CTATCAAGCC	TICIGAAGGI	CATGICAGIG	1850
	CCTCCÄÄTCC	TGTCTGAAGT	TTTCTTTATG	GIAATITTAC	CAAAAGIAAA	1900
10	ÁTCGCTTTGC	TTÅATAÄCCT	TCATTATGCT	CTGACGATTC	TITAGGAATG	1950
	TCAGACATGA	AATAACGCTC	ATCITCTIGA	TCTGGTCGAT	GITTICCAGA	2000
	CAAAAAGICI	TGAAGTTGAA	TGCTACCAGA	TICTGATCIT	CCTCAAACTC	2050
	AAGGICITIG	CCTTGTGTCA	ACAAAGCAAC	AATGCTTTCC	TTAGEGAGCT	2100
	TAACC TTAGA	CATGATGATC	GIAAAAGIIG	TTATAGCTTT	GACCGTATGT	2150
15	AACTCAAGGT	GCGAAAGIGC	AACICIGIAT	CCCGCAGTCG	TTTCTTAGGT	2200
	TCTTAATGTG	ATGATTTGTA	AGACIGAGIG	TTAACGIATG	AACACAAAAT	2250
	TGACACGATT	GCICI 22	65			

The incomplete deduced amino acid sequence of the nonstructural protein gene on TSWV-CPNH1 S RNA is provided below beginning with nucleic acid at position 1 and ending with the nucleic acid codon ending at position 783:

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Gln Val Glu Ser Asn Asn Arg Thr Val Asn Ser Leu Ala Val Lys
                        5
                                                                15
                                           10
     Ser Leu Leu Met Ser Ala Glu Asn Asn Ile Met Pro Asn Ser Gln
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                       20
                                           25
                                                                30
     Ala Ser Thr Asp Ser His Phe Lys Leu Ser Leu Trp Leu Arg Val
     Pro Lys Val Leu Lys Gln Val Ser Ile Gln Lys Leu Phe Lys Val
                                                                60
                       50
                                            55
30
     Ala Gly Asp Glu Thr Asn Lys Thr Phe Tyr Leu Ser Ile Ala Cys
                                                                75
     Ile Pro Asn His Asn Ser Val Glu Thr Ala Leu Asn Ile Thr Val
                       80
                                            85
     Ile Cys Lys His Gln Leu Pro Ile Arg Lys Cys Lys Ala Pro Phe
35
                                           100
                       95
                                                               105
      Glu Leu Ser Met Met Phe Ser Asp Leu Lys Glu Pro Tyr Asn`lle
                                           115
                                                                120
                      110
      Val His Asp Pro Ser Tyr Pro Lys Gly Ser Val Pro Met Leu Trp
                                                                135
                       125
                                           130
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	Leu	Glu	Thr	His	Thr 140	Ser	Leu	His	Lys	Phe 145	Phe	Ala	Thr	Asn	Leu 150
	Gln	Glu	Asp	Val	Ile 155	Ile	Tyr	Thr	Leu	Asn 160	Asn	Leu	Glu	Leu	Thr 165
5	Pro	Gly	Lys	Leu	Asp 170	Leu	Gly	Glu	Arg	Thr 175	Leu	Asn	Tyr	Ser	Glu 180
	Asp	Ala	Tyr	Lys	Arg 185	Asp	Tyr	Phe	Leu	Ser 190	Lys	Thr	Leu	Glu	Cys 195
10	Leu	Pro	Ser	Asn	Thr 200	Gln	Thr	Met	Ser	Tyr 205	Leu	Asp	Ser	Ile	Gln 210
	Ile	Pro	Ser	Trp	Lys 215	Ile	Asp	Phe	Ala	Arg 220	Gly	Glu '	Ile	Lys	Ile 225
	Ser	Pro	Gln	Ser	Ile 230	Ser	Val	Ala	Lys	Ser 235	Leu	Leu	Lys	Leu	Asp 240
15	Leu	Ser	Gly	Ile	Lys 245	Lys	Lys	Glu	Ser	Lys 250		Lys	Glu	Ala	Tyr 255
	Ala	Ser	Gly	Ser	Lys 260	1									

The nucleotide sequence for TSWV-23 depicted below compares 20 closely with the TWSV sequence given above, and contains one-half of the nonstructural gene and one half of the nucleocapsid protein gene. AAATICICIT GCAGTGAAAT CICIGCICAT GITAGCAGAA AACAACATCA 50 TGCCIAACIC TCAAGCITTT GICAAAGCIT CTACIGATIC TAATITICAAG 100 CIGAGCCICT GGCIAAGGGT TCCAAAGGTT TIGAAGCAGA TTTCCATTCA 150 25 GAAATTGTTC AAGGTTGCAG GAGATGAAAC AAATAAAACA TTTTTATTTAT 200 CTATTGCCTG CATTCCAAAC CATAACAGTG TTGAGACAGC TTTAAACATT 250 ACIGITATIT GCAAGCATCA GCICCCAATT CGIAAATGIA AAACTCCTIT 300 TGAATTATCA ATGATGITIT CIGATTIAAA GGAGCCITAC AACATTATIC 350 ATGATCCITC ÁTAICCCCAA AGGATTGITC ATGCICIGCT TGAAACICAC 400 30 ACATCTTTIG CACAAGITCT TIGCAACAAC TIGCAAGAAG AIGIGATCAT 450 CTACACCTIG AACAACCATG AGCTAACTCC TGGAAAGTTA GATTTAGGTG 500 AAATAACITT GAATTACAAT GAAGACGCCT ACAAAAGGAA ATATTICCIT 550 TCAAAAACAC TIGAATGICI TCCATCIAAC ATACAAACIA TGICITATIT 600 AGACAGCATC CAAATCCCTT CCTGGAAGAT AGACTTTGCC AGGGGAGAAA 650 35 TTAAAATTTC TCCACAATCT ATTTCAGTTG CAAAATCTTT GTTAAATCTT 700 GATTTAAGCG GGATTAAAAA GAAAGAATCT AAGATTAAGG AAGCATATGC 750 TTCAGGATCA AAATGATCIT GCTGTGTCCA GCTTTTTCTA ATTATGTTAT 800 GITTATTITC TITCITTACT TATAATTATT TITCIGITIG TCATTICITT 850 CAAATICCIC CIGICIAGIA GAAACCATAA AAACAAAAAT AAAAATAAAA 900

TAAAATCAAA ATAAAATAAA AATCAAAAAA TGAAATAAAA GCAACAAAAA 950 AATTAAAAAA CAAAAAACCA AAAAAGATCC CGAAAGGACA ATTTTGGCCA 1000 ANTIGGGGT TIGITTITGT TTTTGTTTT TITGTTTTT GTTTTTATTT 1050 TTATTITTAT TITTATTITT ATTTTATTIT ATTTTATGIT TTTGTTGTTT 1100 TIGITATITT GITATITATI AAGCACAACA CACAGAAAGCA AACITIAAT 1150 TAAÁCACACT TAITTÁAAAT TTAACACACT AAGCAAGCACA AACAATAAA 1200 GATAAAGAAA GCITTATATA TITATAGGCT TITTIATAAT TIAACITACA 1250 GCTGCTTTTA AGCAAGTTCT GTGAGTTTTG CCTGTTTTTT AACCCCAAAC 1300 ATTICATAGA ACTIGITAAG GGITTCACIG TAATGITCCA TAGCAATACT 1350 10 TOCTITAGCA TIAGGATIGO IGGAGCIAAG TATAGCAGCA TACTOTITOC 1400 CCITCITCAC CIGATCITCA TICATTICAA ATGCTTTICT TITCAGCACA 1450 GIGCAAACIT TICCIAAGGC TICCCIGGIG TCATACITCI TIGGGICGAT 1500 COCGAGATICC TIGIATITIG CATCCIGATA TATAGCCAAG ACAACACIGA 1550 TCATCICAAA GCIATCAACI GAAGCAATAA GAGGTAACCT ACCICCCAGC 1600 15 ATTATEGCAA GCCTCACAGA CTTTGCATCA TCAAGAGGIA ATCCATAGGC 1650 TIGAATCAAA GGGIGGGAAG CAATCTIAGA TITGATAGIA TIGAGATTCT 1700 CAGAATICC 1709

The nucleic acid sequence for TSWV-PCR according to the present invention as depicted below also compares closely with the TSWV 20 sequence given above and covers the whole nucleocapsid protein gene. TTAACACACT AAGCAAGCAC AAACAATAAA GATAAAGAAA GCITTATATA 50 TTIATAGGCT TTTITATAAT TTAACTIACA GCIGCITTIA AGCAAGITCT 100 GIGAGITTIG CCIGITITIT AACCCCAAAC ATTICATAGA ACTIGITAAG 150 GGITTCACTG TAATGTTCCA TAGCAATACT TCCTTTAGCA TTAGGATTGC 200 25 TGGAGCTAAG TATAGCAGCA TACTCTTCC CCTTCTTCAC CTGATCTTCA 250 300 TICATTICAA ATGCTTTTCT TITCAGCACA GIGCAAACIT TICCTAAGGC TICCCIGGIG TCATACTICT TIGGGICGAT CCCGAGATCC TIGIATITIG 350 CATCCTGATA TATAGCCAAG ACAACACTGA TCATCTCAAA GCTATCAACT 400 GAAGCAATAA GAGGTAAGCT ACCICCCAGC ATTATGGCAA GCCTCACAGA 450 30 CTITICCATCA TCAAGAGGIA ATCCATAGGC TTGACTCAAA GGGIGGGAAG 500 CAATCITAGA TITGATAGIA TITGAGATICI CAGAATICCC AGITTCCTCA 550 ACAAGCCTGA CCCTGATCAÁ GCTATCAAGC CTTCTGAAGG TCATGTCAGT 600 GGCICCAATC CIGICIGAAG TITICITIAT GGIAATTITA CCAAAAGIAA 650 AATCGCTTTG CTTAATAACC TTCATTATGC TCTGACGATT CTTCAGGAAT 700 750 35 GICAGACATG AAATAATGCT CATCITITIG ATCIGGICAA GGITTICCAG

ACAÁAAAGIC TIGAÁGITGA ATGCIACCAG ATICIGATCT TCCICAAACT 800 CAAGGICITT GCCITGIGIC AACAAAGCAA CAATGCITTC CITAGIGAGC 850 TIAACCAT 858

Together the cloned TSWV-23 insert overlaps the TSWV-PCR insert, and together they represent the 2028 nucleotides of the TSWV-BL S RNA according to the present invention. This 2028 nucleotide sequence according to the present invention contains a part of the nonstructural gene and whole nucleocapsid protein gene. The combined sequence is:

10	AAATTCTCTT GCAGTGAAAT CTCTGCTCAT GTTAGCAGAA AACAACATCA	50
	TGCCTAACTC TCAAGCTTTT GTCAAAGCTT CTACTGATTC TAATTTCAAG	100
	CIGAGCCICI GGCIAAGGGI TOCAAAGGII TIGAAGCAGA TITCCATICA	150
	GAAATIGITC AAGGITGCAG GAGATGAAAC AAATAAAACA TITITATTIAT	200
	CTATIGCCIG CATICCAAAC CATAACAGIG TIGAGACAGC TITAAACAIT	250
15	ACTGTTATTT GCAAGCATCA GCTCCCAATT CGTAAATGTA AAACTCCTTT	300
	TGAATTATCA ATGATGTTTT CIGATTIAAA GGAGCCTIAC AACATTATTC	350
	ATGATCCTTC ATATCCCCAA AGGATTGTTC ATGCTCTGCT TGAAACTCAC	400
	ACATCITTIG CACAAGITCT TIGCAACAAC TIGCAAGAAG AIGIGAICAT	450
	CTACACCITG AACAACCATG AGCTAACTCC TGGAAAGTTA GATTTAGGTG	500
20	AAATAACITT GAATTACAAT GAAGACGCCT ACAAAAGGAA ATATTICCIT	550
	TCAAAAACAC TIGAATGICI TCCATCIAAC ATACAAACTA TGICITATIT	600
	AGACAGCATC CAAATCCCTT CCTGGAAGAT AGACTTTGCC AGGGGAGAAA	650
	TIAAAATITC TCCACAATCT ATTICAGTIG CAAAATCITT GITAAATCIT	700
	GATTTAAGCG GGATTAAAAA GAAAGAATCT AAGATTAAGG AAGCATATGC	750
25	TICAGGATCA AAATGATCIT GCIGIGICCA GCFFFFTCIA ATTAIGITAT	800
	GTTTATTTIC TTTCTTTACT TATAATTATT TTTCTGTTTG TCATTTCTTT	850
	CAAATTCCIC CIGICIAGIA GAAACCATAA AAACAAAAAT AAAAATAAAA	900
	TAAAATCAAA ATAAAATAAA AATCAAAAAA TGAAATAAAA GCAACAAAAA	950
	AATTAAAAAA CAAAAAAACCA AAAAAGATCC CGAAAGGACA ATTITGGCCA	1000
30	AATTIGGGT TIGITITIGT TITTIGITIT TITGITTITT GITTITATIT	1050
	TTATTTTAT TTTTATTTT ATTTTATTTT ATTTTATGTT TTTGTTGTTT	1100
	TIGITATITI GITATITATI AAGCACAACA CACAGAAAGC AAACITIAAT	1150
	TAAACACACT TATTTAAAAT TTAACACACT AAGCAAGCAC AAACAATAAA	1200
	GATAAAGAAA GCTTTATATA TTTATAGGCT TTTTTATAAT TTAACTTACA	1250
3 5	GCIGCITTIA AGCAAGIICT GIGAGITTIG CCIGITTIIT AACCCCAAAC	1300

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ATTICATAGA ACITGITAAG GGITICACIG TAATGITCCA TAGCAATACT 1350 TOCTTIAGCA TIAGGATIGO IGGAGOTAAG TATAGCAGCA TACTOTTICO 1400 CCITCITCAC CIGATCITCA TICATITICAA AIGCITTICT TITCAGCACA 1450 GIGCAAACIT TICCIAAGGC TICCCIGGIG TCAIACITCI TIGGGICGAT 1500 COCGAGATICC TIGIATITIG CATOCIGATA TATAGOCAAG ACAACACTGA 1550 5 TCATCTCAAA GCTATCAACT GAAGCAATAA GAGGTAAGCT ACCTCCCAGC 1600 1650 ATTATGGCAÁ GOCTCACÁGA CÍTTGCATCA TCAAGAGGTA ATCCATAGGC TIGACICAAA GOGTOGGAAG CAATCITAGA TTIGATAGIA TIGAGATTCT 1700 CAGAATTCCC AGTITCCTCA ACAAGCCIGA CCCTGATCAA GCTATCAAGC 1750 CITCIGAAGG TCATGICAGT GGCICCAATC CIGICIGAAG TITTCITIAT 10 1800 GGEAATTTIA CCAAAAGEAA AATCGCTTIG CITAATAACC TICATTATGC 1850 TCTGACGATT CITCAGGAAT GTCAGACATG AAATAATGCT CATCTTTTTG 1900 ATCIGGICAA GGITTIOCAG ACAAAAAGIC TIGAAGIIGA AIGCIACCAG 1950 ATTCTGATCT TOCTCAAACT CAAGGICTTT GCCTTGTGTC AACAAAGCAA 2000 15 CAATGCTTTC CTTAGTGAGC TTAACCAT 2028

included about 760 bp of the 52 K protein viral component gene, the complete intergenic region (492 bp), and 450 bp of the NP gene (about half of the NP gene). This cloned insert had its 3'-end located exactly at an EcoRI recognition site, which suggested incomplete EcoRI methylation during the cDNA cloning procedure. Although this clone did not contain the complete TSWV-BL NP gene, its sequence was of considerable importance since it had a 450 bp overlap with the sequence of the PCR-engineered NP gene (a total of 2028 bp of the TSWV-BL S RNA is presented in the nucleotide sequence for TSWV). The sequence comparison between this TSWV-BL PCR-engineered and TSWV-CPNH1 NP genes revealed a total of 21 nucleotide differences (2.7%), eight of which encode amino acid replacements (3.1%). Since this PCR engineered NP gene was obtained using Tag polymerase, which is known 30 to incorporate mutations, it is possible that some of these differences were introduced during PCR amplification. However, 15 of these nucleotide differences were located within the overlapping region between the TSWV-BL cDNA and PCR clones, and all but one of these nucleotide differences (position 1702 of TSWV; position 485 of TSWV-

This comparison showed that cDNA insert of clone pTSWVS-23

35 PCR)) are shared by both TSWV-BL S RNA derived clones. This

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comparison clearly showed that the PCR amplification did not contribute greatly, if at all, to the difference between the nucleotide sequences of these two cloned NP gene regions. The nucleotide difference at position 1702 resulted in the amino acid replacement of the with Ser, and even this difference could be due to the lack of homogeneity within the TSWV-BL isolate.

EXAMPLE V

Agrobacterium-mediated transformation:

Leaf discs of Nicotiāna tabacum var Havana cv 423 were inoculated with the Agrobacterium strain LBA4404 (ClonTech) 10 containing the vector pBIN19-NP+ or the control plasmid pBIN19, by soaking overnight in a liquid culture of the Agobacterium, and the inoculated leaf discs were incubated on non-selective MS medium for 3 days. [see Science 227:1229 (1985)]. Transformed cells were selected 15 and regenerated in MS medium containing 300 µg/ml kanamycin and 500 µg/ml carbenicillin for shoot regeneration. Roots were induced after transfer of plantlets to hormone-free medium. Rooted transformants were transferred to soil and grown under greenhouse conditions. The MS medium contains full strength MS salt (Sigma), 30 g/l sucrose, 1 mg/l 20 BA and 1 ml of B5 vitamins [1 mg/ml Nicotinic acid, 10 mg/ml Thiamine (HCI), 1 mg/ml Pyridoxine (HCI), 100 mg/ml Myo-Inositol]. Transgenic plants were self-pollinated and seeds were selectively germinated on kanamycin medium.

EXAMPLE VI

25 Serological detection of proteins:

Double antibody sandwich enzyme-linked immunosorbent assay (DAS-ELISA) was used to detect the expression of NP gene in transgenic plants with polyclonal antibodies against the TSWV-BL NP. Each sample was prepared by grinding a leaf disc (about 0.05 g) from the top second leaf of the plant in 3 ml of an enzyme conjugate buffer [phosphate-buffered saline, 0.05% Tween 20, 2% polyvinylpyrrolidone 40, and 0.2% ovalbumin]. For tobacco protoplasts, the cell extracts after centrifugation were directly used for the assay. A ten- and three-fold dilutions of the samples from both transgenic plants and tobacco protoplasts were made just before DAS-ELISA.

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For Western blots, a leaf disc (about 0.05 g) was ground in 0.25 ml of 2X SDS/sample buffer (0.126 M Tris buffer, 20% glycerol, 2% SDS, 2% 2-mercaptoethanol, and 0.01 mg/ml bromphenol blue). The homogenates were centrifuged and the supernatants were boiled before loading. Proteins (10-20 μl sample/lane) were separated and blotted onto a membrane. The membrane was then processed following the manufacturer's immunoselect kit instruction manual (Gibco BRL Life Technologies Inc.). Antibodies to the whole virion were preabsorbed with cell extracts from health tobacco plants [See Plant Disease 70:501 (1986)], and were used in Western blot at a concentration of 2 μg/ml.

Serological reactions of TSWV isolates (TSWV-BL, Arkansas, 10W pakchoy, Begonia or Brazil) were assayed in DAS-ELISA using antibodies raised against TSWV-BL virion, or the NP of TSWV-BL or TSWV-I.

EXAMPLE VII

Inoculation of transgenic plants with TSWV isolates.

Inocula were prepared by infecting *Nicotiana benthamiana* Domin. with different TSWV isolates and grinding infected leaves (0.5 g) of *N. benthamiana* plants (1 to 2 weeks after inoculation) in 15 ml. of a buffer (0.033 M KH₂PO₄, 0.067 M K₂HPO₄ and 0.01 M Na₂SO₃). The inoculum extracts were immediately rubbed on corundum-dusted leaves of transgenic plants and the inoculated leaves were subsequently rinsed with H₂O. Because TSWV is highly unstable in vitro after grinding, each batch of inoculum was used to first inoculate NP(+) plants containing the NP gene; the last inoculated plants of each inoculum were always control NP(-) plants containing the vector sequence alone to assure that a particular virus inoculum was still infective at the end of inoculation.

Data on local lesions and systemic infections were taken 7-15 days after inoculation and expressed in the following table as the number of plants systemically infected over the number of plants inoculated, except where indicated. In this table, the data collected under "ELISA" is the data of R₀ lines from which the R₁ plants were derived; the Begonia isolate induced local lesions on the R₁ plants, and the resistance was expressed as the number of plants producing local

lesions over the number of plants inoculated; and NT indicates that there was no test.

Reactions of R1 plants expressing the nucleocapsid protein (NP) gene of tomato spotted wilt virus (TSWV) to inoculation with TSWV isolates.

5		Reactions to TSWV isolates					
		ELIŜA: (R0 pl.)	BL	Arkansas	10W Pakchoy	Begonia	Brazil
	Ro line			•			,
10	NP(+)2	0.015	0/20	4/25	3/24	29/40	36/36
	NP(+)4	0.386	6/30	21/23	18/21	', 9/48	42/42
	NP(+)9	0.327	0/20	NT	20/20	_	_
	NP(+)14	0.040	0/20	_	9/20	8/18	18/18
	NP(+)21	0.042	0/15	5/15	3/15	2/4	6/6
15	NP(+)22	0.142	0/20		15/20	31/36	36/36
	NP(+)23	0.317	0/20		16/20		
-	NP(-)	•	42/42	24/24	62/62	66/66	54/54

resides in the S RNA component of TSWV, was approached using two 20 strategies. The cDNA cloning strategy yielded several clones containing cDNA inserts derived from TSWV-BL S RNA, as identified by hybridization against an oligomer probe complementary to the TSWV-CPNH1 S RNA. Clone pTSWVS-23 contained the longest insert, which mapped at about 1.7 kb in length. The second strategy utilized the 25 published sequence of TSWV-CPNH1 S RNA and PCR to amplify and engineer the NP gene for expression directly from total TSWV-BL RNA. Oligomer primers JLS90-46 and -47 were synthesized, with JLS90-46 being complementary to the S RNA in the 5'-coding region of the NP gene (positions 2051-2073 of the TSWV-CPNH1) while JLS90-47 being of 30 the 3'-noncoding region of the NP gene (positions 1218 to 1237 of the TSWV-CPNH1). Both of the primers contain the recognition site for the restriction enzyme Ncol for subsequent cloning, and the primer JLS90-46 has a plant consensus translation initiation codon sequence (AAXXATGG), which upon amplification was expected to fuse the 35 translation initiation codon to the third codon (GTT) of the NP gene. Fusion of the translation initiation codon to the third codon of the

As described above, the isolation of the TSWV-BL NP gene, which

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TSWV-BL NP gene was done to preserve the *Ncol* recognition site while not incorporating any amino acid codons. Thus, expression of the PCR-engineered TSWV NP gene would yield a TSWV-BL NP that was two amino acids (Ser-Lys) shorter at the N-terminus than the native NP.

This specifically-amplified DNA fragment, of about 850 bp, was digested with Ncol and cloned into the plant expression vector pB1525. The orientation of the TSWV-BL NP gene with respect to the CaMV 35S promoter was determined by restriction enzyme site mapping (EcoRI, HindIII, Avai and AlwNI). Several clones were isolated that contain the insert in the proper orientation (pB1525-NP+) and others that contain the insert in the opposite orientation (pB1525-NP-). This restriction enzyme site mapping data also showed that the inserts of clones pB1525-NP+ contained restriction enzyme sites that were identical to those found in the TSWV-CPNH1 NP gene. The expression of TSWV-BL NP gene was thus controlled by a double CaMV 35S promoter fused to the 5'-untranslated leader sequence of alfalfa mosaic virus (ALMV) of the expression vector pB1525. Expression vectors that utilize the stacked double CaMV 35S promoter elements yield higher levels of mRNA transcription than similar vectors that utilize a single 35S promoter element.

Three pB1525-NP+clones were translently expressed in tobacco protoplasts to confirm that the amplified DNA fragment encoded the NP. To achieve this, the clones were transferred into tobacco protoplasts by the PEG method, and after two days of incubation the expressed NP was detected by DAS-ELISA using antibodies against the whole TSWV-BL virion. High levels of NP were produced in tobacco protoplasts harboring the NP gene in plasmid pB1525-NP+; while no NP was detected in tobacco protoplasts transformed with the antisense NP sequence (pB1525-NP-).

As described previously, the PCR-engineered insert of clone pBI525-NP+ and teh cDNA insert of the clone pTSWV-23 were subjected to double stranded sequencing. The sequence analysis of the cDNA and the PCR clones revealed inserts of 1.71 kb and 865 bp, respectively which, when compared with the sequence TSWV-CPNH1 S RNA, shows that cDNA insert of clone pTSWV-23 includes about 760 bp of the 52 K

protein viral component gene, the complete intergenic region (492 bp), and 450 bp of the NP gene (about one-half of the gene). This cloned insert has its 3'-end located exactly at an *EcoRI* recognition site suggesting incomplete *EcoRI* methylation during the cDNA cloning

- procedure. Although this clone does not contain the complete TSWV-BL NP gene, its sequence is of considerable importance since it has a 450 bp overlap with the sequence of the PCR-engineered NP gene. The sequence comparison between this TSWV-BL PCR-engineered and TSWV-CPNH1 NP genes reveals a total of 21 nucleotide differences (2.7%),
- eight of which encode amino acid replacements (3.1%). Since this PCR-engineered NP gene was obtained using *Taq* polymerase, which is known to incorporate mutations, it is possible that some of these differences were introduced during PCR amplification. However, 15 of these nucleotide differences are located within the overlapping region
- between the TSWV-BL cDNA and PCR clones, and all but one of these differences (position 1702) are present in both TSWV-BL S RNA derived clones. This comparison clearly shows that the PCR amplification did not contribute greatly, if at all, to the difference between the nucleotide sequences of these two NP genes. The nucleotide difference
- at position 1702 results in the amino acid replacement of lie with Ser, and even this difference could be due to the lack of homogeneity within the TSWV-BL isolate.

The possibility that the nucleotide differences can be attributed to divergence among the TSWV isolates is also supported by

25 comparisons with other sequenced regions among TSWV-CPNH1, TSWV-L3, and TSWV-BI S RNAs. These comparisons are tabulated below:

Percent nucleotide and amino acid sequence differences for the comparison of TSWV S RNA component from isolates CPNH1, L3 and BL^a

		52 K Protei	n Gene	Intergenic	NP Gene	<u>.</u>
30	Comparison	Nucleotide #	Amino Acid	Nucleotide	Nucleotide	Amino Acid
	CPNH1/L3	68/1396 ^b (4.9) ^c	49/464(10.6)	46/511(9.0)	24/777(3.1)	4/258(1.6)
	CPNH1/BL	21/758(4.1)	23/251(9.2)	26/496(5.2)	19/765(2.5)	8/255(3.1)
	L3/BL	38/765(5.0)	20/254(7.9)	38/498(7.6)	19/767(2.5)	4/255(1.6)

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Comparisons are made using the sequence information available from the particular component region of TSWV-BL. The comparison for the TSWV-BL NP gene includes the combined sequence information from the cDNA clone, pTSWVS-23 and PCR-engineered insert.

Comparison numbers are total differences (nucleotides or amino acids) divided by total number of positions (nucleotides or amino acids) compared. For both nucleotide and amino acid calculation gaps, regardless of length, were counted as one mismatch

Numbers in parenthèses are percentages.

The nucleotide sequence of the NP genes from the CPNH1 and L3 isolates differ from each other by 3.1% and from the BL isolate by 'nearly a similar degree (2.5%). However, the NP amino acid sequences between CPNH1 and BL isolates differ by a considerably larger amount than they differ between the L3 and BL or CPNH1 and L3 isolates. The results tabulated above also reveal that the NP gene region of these TSWV isolates is subject to a higher degree of selective pressure than the 52 K protein as the differences among the amino acid sequences of the 52 K protein range between 7.9 to 10.6%, more than twice that found for the amino acid sequence of the NPs. Nucleotide sequence divergence is highest among the intergenic regions, indicating that this region is subject to less selective pressure than either genetic region.

The presence of NP gene sequences in transgenic plants was first confirmed by PCR analysis. A NP DNA fragment of about 800 bp was specifically amplified from the total DNAs of transgenic NP(+) plants using the primers homologous to sequences flanking the NP gene, whereas no corresponding fragment was detected in control NP(-) plants. Expression of the NP gene was assayed in each R0 transgenic plant by DAS-ELISA, and the results are presented in the following table:

Reactions of R0 transgenic plants expressing the nucleocapsid protein (NP) gene of tomato spotted wilt virus (TSWV) to inoculation with TSWV-BL isolate

	plant age	R ₀ clone	ELISAa	Lesions/leaf ^b	NP(+):NP(-)C
	7-8 leaves:				
5		NP(+)1	0.374	7 (199)	1:28
	X.	NP(+)2	0.015	0 (199)	0:199
		NP(+)3	0.407	23 (102)	1:4
		NP(+)4	0.386	2 (102)	1:51
		NP(+)5	0.023	0 (124)	0:124
10		NP(+)6	0.197	35 (325)	1:9
		NP(+)7	0.124	1 (325)	1:325
	9-10 leaves:				
		NP(+)8	0.344	36 (36)	1:1
		NP(+)9	0.327	2 (20)	1:10
15		NP(+)10	0.406	34 (33)	1:1
		NP(+)11	0.156	5 (20)	1:4
		NP(+)12	0.133	9 (57)	1:6
4		NP(+)13	0.144	2 (7)	1:4
		NP(+)14	0.040	0 (19)	0:19
20		NP(+)16	0.053	0 (10)	0:10
	5-6 leaves:				
		NP(+)20	0.487	203 (117)	2:1
		NP(+)21	0.042	0 (117)	0:117
		NP(+)22	0.142	0 (208)	0:208
25		NP(+)23	0.317	223 (208)	1:1
		NP(+)24	0.051	0 (35)	0:35
		NP(+)25	0.286	13 (35)	1:3
		NP(+)26	0.037	0 (22)	0:22
		NP(+)27	0.425	305 (22)	14:1

- aproduction of the NP in transgenic plants was assayed by double antibody sandwich enzyme-linked immunosorbent assay (DAS-ELISA); concentration of antibodies against viron for coating: 1 μg/ml; dilution of conjugate to the NP of TSWV-BL: 1:250; results taken 150 min. after adding substrate; readings at 405 nm.
- 35 blocal lesions that developed on inoculated leaves were counted 7 days after inoculation. Data represent the average of three inoculated leaves. Data in parentheses are the number of lesions produced from control NP(-) plants inoculated with the same inoculum.

Cthe ratio of local lesions that developed on NP(+) plants transformed with pBIN19-

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NP+ versus local lesions that developed on the control NP(-) plant when inoculated with the same inoculum.

Of the 23 NP(+) clones, 10 produced high levels of NP, 5 accumulated intermediate levels of NP, and the remaining 8 produced low levels of NP. The size of the NP expressed in transgenic plants was analyzed using Western blot. Many polypeptides from tobacco extracts were reactive to the antibodies against the whole viron even though the antibodies were pre-absorbed with extracts from healthy tobacco plants. Of those, only one band was unique to the pattern of polypeptides from tobacco plants transformed with the NP gene. This polypeptide was estimated to be around 29 kDa, which is near the expected size of the native NP. No antibody reactive-protein band of similar size was found in extracts from transgenic plants containing the vector pBIN19.

Inoculation of tobacco leaves with TSWV-BL isolate could result in either systemic infection or necrotic local lesions, depending upon weather conditions and physiological stages of plants. When Ro plants were tested with TSWV-BL for viral resistance, TSWV-BL induced typical necrotic lesions on the inoculated leaves of control NP(-) plants 6-8 days after inoculation. However, transgenic NP(+) plants showed a spectrum of resistance to the virus when compared to control NP(-) plants. Eleven of the 23 NP(+) plants did not develop any local lesion or the number of lesions that developed was at least 20-fold less than that on the corresponding inoculated NP(-) plants. Three NP(+) plants had intermediate reactions (5- to 19-fold less lesions than controls) while the remaining 9 plants had low or no resistance. None of the inoculated NP(+) or NP(-) plants showed systemic infection. symptomless Ro plants were monitored until the end of their life cycle, and no symptom was observed throughout their life cycles. The inoculated leaves of the symptomless NP(+) plants were checked for the presence of the virus on the leaves of C, quinoa plants. No virus was recovered from TSWV-BL-challenged leaves of highly resistant NP(+) plants, suggesting that the virus cold not replicate or spread in these NP(+) plants.

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Leaf discs from selected R_0 plants were subcloned, and the regenerated plantlets were challenged by the virus. All subcloned R_0 plants displayed levels of resistance similar to their corresponding original R_0 plants.

Since TSWV is widespread and many biologically distant strains exist, the effectiveness of the transgenic plants to resist infections by different TSWV isolates were also tested. Five TSWV isolates were chosen in this study to challenge R1 plants germinated on kanamycincontaining medium: TSWV-BL, Arkansas, 10W pakchoy, Begonia and Brazil. The first three isolates were reactive to the antibodies against the whole virion and the NP of TSWV-BL (the common TSWV "L" serogroup) (see figure 5). Begonia isolate reacted strongly to the antibodies against the NP of TSWV-I (the "I" serogroup) but not to those raised against the TSWV-BL NP, and therefore belonged to the "I" serodroup. No detectable reaction of Brazil isolate was found to the antibodies against either the NP of the TSWV-BL or the TSWV-I serogroup, and it was weakly reactive to the antibodies against the whole viron of TSWV-BL. Moreover, this isolate caused systemic mottle and crinkle on the leaves of infected tobacco and N. benthamiana, but did not infect squash or cucumbers indicating that it is a distinct isolate from the cucurbit isolate. These results indicate that this isolate may be considered to be a third serogroup.

Seedlings derived from seven R₀ lines were germinated on kanamycin medium and inoculated with the above TSWV isolates.

- Infectivity data were recorded daily starting seven days after inoculation. Plants inoculated with TSWV-BL, Arkansas, 10W pakehoy or Brazil isolates were rated susceptible if virus symptoms were observed on uninoculated leaves. Plants inoculated with the Begonia isolate were rated susceptible if local lesions were observed on inoculated leaves, since this isolate does not cause systemic infection
- in tobacco. All inoculated control NP(-) R₁ plants were susceptible to infection by these five isolates. They were systemically infected 12 days after inoculation except that transgenic R₁ plants inoculated with Begonia produced only local lesions on the inoculated leaves. However,
- 35 almost all NP(+) R₁ plants were highly resistant to the homologous

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isolate TSWV-BL, while much lower percentages of NP(+) R₁ plants were resistant to heterologous isolates Arkansas, 10W pakchoy and Begonia. On the other hand, all NP(+) R₁ plants from the seven transgenic lines were susceptible to the Brazil Isolate, even though a slight delay (1 to 2 days) in symptom expression was observed in some of the high NP-expressing NP(+) R₁ plants from line NP(+)4.

Resistant R1 plants remained symptomless throughout their life cycles. The inoculated leaves of seventeen symptom less NP(+) plants were checked for the presence of the virus by back inoculation on leaves of *Chenopodium quinoa* plants. No virus was recovered from the inoculated leaves of symptomless NP(+) plants, suggesting that the virus could not replicate or spread in these NP(+) plants.

The relationship between the level of NP accumulation in transgenic plants and the degree of resistance to heterologous TSWV isolates was also studied. Analysis of the data described above suggested that R₁ plants derived from R₀ lines with low levels of NP offered the best resistance to the heterologous isolates of the "L" serogroup (Arkansas and 10W pakchoy) while R₁ from a R₀ line with high level of NP were resistant to the Begonia isolate, which belongs to the "I" serogroup. For example, an average 76% of inoculated R₁ plants from low NP expressing lines NP(+) 2, 14, and 21 were resistant to infections by the Arkansas and 10W pakchoy isolates, while resistance to these isolates was observed in only 11% of similarly inoculated plants from high NP expressing lines NP(+)4, 9, and 23. On the other hand, the Begonia isolate infected 79% of R₁ plants from the low NP expressing line NP(+)2, 14, and 21 but only 19% from high NP expressing line NP(+)4.

Therefore, it was concluded that the transgenic R₁ plants expressing low levels of the NP gene were highly resistant to infection with the isolate 10W pakehoy (the "L" serogroup), but not to Begonia isolate (the "I" serogroup). In contrast, the highly NP-expressing R₁ plants were very resistant to infection by Begonia isolate but not to infection by the isolate from 10W pakehoy.

Thus, it was of interest to accurately quantitate the relation of NP expression in individual plants with resistance to the heterologous

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isolates. In a number of inoculation experiments reported herein, leaf samples of transgenic plants were taken before inoculating with the Arkansas and 10W pakehov isolates. Samples were also taken from non-inoculated leaves of plants inoculated with the Begonia isolate after observations of the apparent relation between NP expression levels and resistance were made. The latter method of sampling could be done without interferee from infection by the Begonia isolate because this isolate does not cause systemic infection in tobacco nor reacts with antibodies to the TSWV-BL NP. All samples were assayed for relative NP levels by DAS-ELISA using antibodies raised to isolated NP of TSWV-BL. Figures 5 and 6 show the relation between NP levels in transgenic R₁ plants (irrespective of the R₀ lines they came from) and their resistance to the Arkansas and 10W pakchov isolates or to the Begonia isolate. Nearly all transgenic R1 plants with very low or undetectable ELISA reactions (0-0.05 OD405nm) were resistant to infections by the Arkansas and 10W pakchoy isolates (the "L" serogroup) but susceptible to the Begonia isolate (the "I" serogroup). In contrast, almost all R₁ plants that gave high ELISA reactions (0.4-1.0 OD405nm) were resistant to the Begonia isolate but susceptible to the Arkansas and 10W pakehov isolates.

The double-stranded (ds) RNA was isolated from the *N*. benthamiana plants infected with TSWV-B using a combination of methods [See Acta Horticulturae 186:51 (1986), and Can. Plant Dis Surv 68:93(1988)] which have been successfully used for isolation of dsRNA from tissue infected with grapevine leafroll virus. The dsRNA was chosen for the cDNA synthesis since isolation of the virus particle from this isolate has not been possible [see Plant Disease 74:154 (1990)]. In order to make a cDNA library specific to the S RNA of TSWV-B, the double stranded S RNA was gel-purified, denatured by methyl-mercury treatment, and subjected to cDNA synthesis procedure provided by Promega using random primers. The synthesized cDNA fragments were cloned via an EcoRI adaptor into the *Eco*RI digested λ ZAPII (Strategene), and positive clones were identified by colony hybridization using the cDNA probes prepared by reverse transcription of gel-purified S RNA. Dozens of positive clones were analyzed on

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agarose gels and only three overlapping clones containing the largest inserts (L1, L22 and L30) were selected (see figure 3), covering nearly entire TSWV-B S RNA.

The nucleotide sequences of the inserts in clones L1, L22 and L30 were determined from both DNA strands, first by the universal and reverse primers and then by the internal primers designed for sequencing the S RNA of TSWV-B. Sequencing was done using the Sanger dideoxyribohucleotide method, T7 polymerase (U.S. Biochemicals, Sequenase TM), and the double-stranded sequencing procedure described by Siemieniak [see Analyt. Biochem. 192:441 (1991)]. The sequence analyses of these clones revealed inserts of 1.994 kb, 2.368 kb and 1.576 kb, respectively, and these sequences represented 93% of the S RNA genome (see figure 3). The assembled sequence was analyzed by comparisons with sequences of TSWV isolates CONH1, L3, I, and BL using computer programs available from the Genetics Computer Group (GCG, Madison, WI).

Computer analysis showed that the assembled sequence of 2.842 kb covered the complete 52 K nonstructural protein gene, the complete intergenic region (629 bp), and 737 bp of the NP gene (only 39 N-20 terminal nucleotides of the N gene were not represented). In order to obtain this missing region of the N gene, a primer TTCTGGTCTTCTTCAAACTCA, identical to a sequence 62 nucleotides from the initiation codon of the N gene, was end-labeled with polynucleotide kinase to screen the cDNA library described above. Five putative clones were obtained. Sequence analysis of the five clones showed that only clones S6 and S7 contain these 39 missing nucleotides of the N gene. The latter clone also included the extreme 3' end of the S RNA.

The 5' extreme end of the S RNA was obtained using the 5' RACE System (GIBCO). Both ssRNA of TSWV-B and total RNAs isolated from tobacco plants infected with TSWV-B were used to synthesize first strand cDNA with an oligonucleotide (5'-CTGTAGCCATGAGCAAAG) complementary to the nucleotide positons 746-763 of te TSWV-B S RNA. The 3'-end of the first strand cDNA was tailed with dCTP using terminal deoxynucleotidyl transferase. Tailed cDNA was then amplified

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by PCR using an anshor primer that anneals to the homopolymeric tail, and an oligonucleotide (5'-TTATATCTTCTTCTTGGA) that anneals to the nucleotide positions 512-529 of the TSWV-B S RNA. The PCR-amplified fragement was gel-purified and directly cloned into the T-vector pT7Blue (Novagen) for sequence analysis. Eight independent clones were sequenced with an oligomer primer (5'-GTTCTGAGATTTGCTAGT) close to the 5' region of the S RNA (nucleotide positions 40-57 of the TSWV-B S RNA). Six of the resulting clones contained the 5' extreme end of the S RNA and the 5'-terminal nucleotide sequence from these clones was identical. Thus, the complete nucleotide sequence of the TSWB-B S RNA is 3049 nucleotides in length.

Thus these two clones together with the three clones previously sequenced (L1, L22, L30, S6 and S7) covered a total of 3032 nucleotides depicted above. Comparisons with the terminal sequences of TSWV-CPNH1 and TSWV-I revealed that although the extreme 5' end of 18 nucleotides was not represented in the assembled sequence, the extreme 3'-terminus of the TSWV-B S RNA is identical to the extreme 3' end of the TSWV-I S RNA and is only one out of fifteen nucleotides different from the extreme 3' end of TSWV-CPNH1. The conservation of the terminal sequence among TSWV isolates is consistent with observations of the other members of *Bunyaviridae* genera, and supports the hypothesis that the terminal sequences might form stable base-paired structure, which could be involved in its replication and encapsulation.

The complete nucleotide sequence of the S RNA genome of TSWV-B (the Brazilian isolate discussed above) according to the present invention is:

-	AGAGCAATTG GGTCATTTT TATTCTAAAT CGAACCTCAA CTAGCAA	AATC 50
30	TCAGAACTGT AATAAGCACA AGAGCACAAG AGCCACAATG TCATCAG	GIG 100
	TITATGAATC GATCATTCAG ACAAAGGCIT CAGITTGGGG ATCGAC	AGCA 150
	TCTGGTAAGT CCATCGTGGA TTCTTACTGG ATTTATGAGT TTCCAA	CIGG 200
	TTCTCCACTG GTTCAAACTC AGTTGTACTC TGATTCGAGG AGCAAA	AGTA 250
	GCTTOGGCTA CACTTOAAAA ATTGGTGATA TTOCTGCTGT AGAGGA	GGAA 300
35	ATTTIATCIC AGAACGITCA TATCCCAGIG TITGATGATA TIGATT	TCAG 350

CATCAATATC AATGATTCIT TCTTGGCAAT TTCTGTTTGT TCCAACACAG 400 450 TTAACACCAA TGGAGTGAAG GATCAGGGTC ATCITAAAGT TCTTTCTCTT 500 GCCCAATIGC ATCCCITIGA ACCIGIGATG AGCAGGICAG AGATIGCTAG 550 CAGATTICOGO CITOCAAGÀAG ÁÁGATATAÁT TOCTGATGAC ÁÁATATATAT 600 CICCICCIAA CAACCCATICI CICICCICIG TCAAACAACA TACTTACAAA 5 650 GTOCANATICA COCACAATICA COCTTTAGGC AAAGTGAATG TTCTTTCTCC TAACACAAAT GITCATGAGT GOCTGTATAG TITCAAACCA AATTICAACC 700 AGATOGAAAG TAATAAGAGA ACTGTAAATT CTCTTGCAGT CAAATCTTTG 750 800 CICATGCTTA CAGAAACAA CATTATGCCT AACTCTCAAG CTTTTGTTAA AGCITICTACT GATTCTCATT TTAAGITGAG CCITTGGCTG AGAATTCCAA 850 10 900 AAGTTTTGAA GCAAATAGOC ATACAGAAGC TCTTCAAGTT TGCAGGAGAC 950 GAAACCGGTA AAAGTTTGTA TTTGTCTATT GCATGCATCC CAAATCACAA 1000 CAGTGTGGAA ÁCAGCTTTAA ATGTCACTGT TATATGTAGA CATCAGCTTC 1050 CAATCCCTAA GTOCAAAGCT CCTTTTGAAT TATCAATGAT TTTCTCCGAT 15 CTGAAAGAGC CTTACAAGAC TGTGCATGAT CCTTCATATC CTCAAAGGAT 1100 TGITCATGCT TTGCTTGAGA CICACACITC CTTTGCACAA GITCICIGCA 1150 1200 ACAAGCIGCA AGAAGATGIG ATCATATATA CTATAAACAG CCCTGAACTA 1250 ACCCCAGCTA AGCTGGATCT AGGTGAAAGA ACCTTGAACT ACAGTGAAGA 1300 TECTTOGAAG AAGAAGTATT TTCTTTCAAA AACACTOGAA TECTTECCAG 1350 20 TAAATGIGCA GACTATGICT TATTIGGATA GCATCCAGAT TCCTICATGG 1400 AAGATAGACT TTGCCAGAGG AGAGATCAGA ATCTCCCCTC AATCTACTCC 1450 TATTCCAAGA TCTTTCCTCA AGCTGGATTT GAGCAAGATC AAGGAAAAGA AGICCITGAC TIGGGAAACA TCCAGCIAIG ATCIAGAATA AAAGIGGCIC 1500 ATACTACICT AAGTAGTATT TGICAACITG CITATCCTTT ATGITGTTTA 1550 TITCITITAA ATCTAAAGTA AGITAGATIC AAGTAGITTA GIATCCTATA 1600 25 GCATTATTAC AAAAAATACA AAAAAATACA AAAAAATACA AAAAATATAA 1650 AAAACCCAAA AAGATCCCAA ÄAGGGACGAT TIGGIIGATT TACICIGITT 1700 TAGGCTTATC TAAGCTGCTT TTGTTTGAGC AAAATAACAT TGTAACATGC 1750 AATAACIGGA ATTTAAAGIC CIAAAAGAG TITCAAAGGA CAGCITAGCC 1800 AAAATIGGIT TITGIITIIG TTITITIGIT TITIGITITT TIGITITATT 1850 30 1900 TITATITITA GITTATITIT TGITTITGIT ATTITATIT TTATITITATT 1950 TICTITIATT TIATTIATAT ATATATCAAA CACAATCCAC ACAAATAATT TIAATTICAA ACATTCIACI GATTIAACAC ACITAGCCIG ACITTATCAC 2000 ACTTAACACG CITAGITAGG CITTAACACA CIGAACTGAA TTAAAACACA 2050 CITAGIATTA TGCATCICIT AATTAACACA CITTAATAAT ATGCATCICT 35 2100

	GAATCAGCCT TAAAGAAGCT	TTTATGCAAC	ACCAGCAATC TIGGCCICIT 2	2150
	TCITAACICC AAACATTICA	TAGAATTIGT	CAAGATTATC ACIGTAATAG	2 20 0
	TOCATAGCAA TGCTTCCCTT	AGCATTGGGA	TTGCAAGAAC TAAGTATCTT	2250
•	GGCATATICT TICCCITIGI	TTATCIGIGC	ATCATCCATT GIAAATCCTT	2300
5	TGCTTTTÄAG CACTGTGCÄA	ÁCCITCCCCA	GAGCITCCIT AGIGITGIAC	2350
	TRACTIGGIT CAATCCCTAA	CICCITGIAC	TTTGCATCTT GATATATGGC	2400
	AAGAACAACA CIGATCATCT	CGAAGCIGIC	AACAGAAGCA ATGAGAGGGA	2450
	TACTACCICC AAGCATTATA	GCAAGTCTCA	CAGATITICC ATCTCCCAGA	2500
•	GGCAGCCCGT AAGCTTGGAC	CAAAGGGIGG	GAGGCAATIT TIGCTITGAT	255Ó
10	AATAGCAAGA TICICATIGI	TIGCAGICIC	TICIATGAGC TICACICTIA	2600
	TCATGCIATC AAGCCTCCTG	AAAGICATAT	CCTTAGCTCC AACTCTTTCA	2650
	GAATTTTICT TTATCGTGAC	CITACCAAAA	GIAAAATCAC TTIGGITCAC	2700
	AACTITCATA ATGCCTTGGC	GATTCITCAA	GAAAGICAAA CATGAAGIGA	2750
	TACTCATTTT CTTAATCAGG	TCAAGATTIT	CCIGACAGAA AGICITAAAG	2800
15				2850
			•	2900
				2950
	•			3000
4.0				3049
20			ences of the nonstructural	
		nucleocapsid	proteins according to the	present
	invention are:			
	Met Ser Ser Gly Val 1	lyr Glu Ser i	Ile Ile Gln Thr Lys Ala 10	Ser 15
25	_	Ala Ser Gly :	Lys Ser Ile Val Asp Ser	
	20		25	30
	Trp He Tyr Glu Phe F	Pro Thr Gly	Ser Pro Leu Val Gln Thr	
		Arg Ser Lvs	40 Ser Ser Phe Gly Tyr Thr	45 Ser
30	50		5 5	60
		Pro Ala Val	Glu Glu Ile Leu Ser	
	65 Asn Val His Ile Pro V	Val Phe Aso	70 Asp Ile Asp Phe Ser Ile	75
	80	·	85	90
3 5		Leu Ala Ile	Ser Val Cys Ser Asn Thr	`
	95 Asn Thr Asn Glv Val	lvs His Gln	100 Gly His Leu Lys Val Leu	105 Ser
	110	are the out	115	120

	Leu .	Ala	Gln	Leu	His 125	Pro	Phe	Glu	Pro	Val 130	Met	Ser	Arg		Glu 135
	Ile .	Ala	Ser	Arg		Arg	Leu	Gln	Glu		Asp	Ile	Ile		
5	Åsp	Lys	Tyr	Ile		Ala	Ala	Asn	Lys		Ser	Leu	Ser	Cys	Val 165
	Lys	Glu	His	Thr	Tyr 170	Lys	Val	Glu	Met	Ser 175	His	Asn	Gln	Ala	Leu 180
10	-	_			185					190			His		195
					200	•				205		1			Asn ' 210
					215				•	220			Met		225
1 5	Glu	Asn	Asn	Ile	Met 230		Asn	Ser	Gln	Ala 235	Phe	Val	Lys	Ala	Ser 240
	Thr	Asp	Ser	His	Phe 245		Leu	Ser	Leu	Gln 250		Arg	Ile	Pro	Lys 255
20	Val	Leu	Lys	Gln	Ile 260		Ile	Gln	Lys	Leu 265		Lys	Phe	Ala	Gly 270
77	Asp	Glu	Thr	Gly		Ser	Phe	Yyr	Let	Ser 280	Ile	Ala	Cys	Ile	Pro 285
	Asn	His	s Asn	Ser		. Glu	Thi	: Ala	Let	295		Thr	<u>Val</u>	Ile	Cys 300
25	Arg	His	s Glr	ı Lev		ıle	Pro	Lys	Ser	: Lys 310		Pro	Phe	Glu	Le u 31 5
	Ser	Met	: Ile	e Phe		: Asp	Let	ı Lys	Gl:	2 Pro 325		: Asn	Thr	Val	His 330
30	Asp	Pro	o Sei	с Туз	2 Pro 33!		ı Ar	g Ile	e Va	1 His 340		Leu	i Leu	Glu	Thr 345
	His	Th	r Sei	r Phe		a Gli	n Va	l Lei	ı Cy	s Asi 355		s Leu	ı Glr	Glu	1 Asp 360
	Val	. Ile	e Ile	e Ty:		r Ile	e As	n Sei	r Pr		ı Le	ı Thi	r Pro	Ala	a Lys 375
3 5	Leu	ı As	p Le	u Gl		u Ar	g Th	r Le	u <u>As</u>	n Ty 38		r Gli	j Asp	Ala	390
	Lys	s Ly	s Ly	s Ty		e Le	u Se	r Ly	s-Th		u Gl	u Cy:	s Lei	ı Pro	o Val 405
40	Ası	n Va	l Gl	n Th		t Se	r Ty	r Le	u As		r Il	e Gl	n Ile	e Pr	o Ser 420
40	Tr	o Ly	s Il	e As		e Al	a Ar	g Gl	y Gl		e Ar	g Il	e Se	r Pr	o Gln 435
	Se:	r Th	ır Pr	ro Il		a Ar	g Se	er Le	u Le		s Le	u As	p Le	u Se	r Lys 450

Ile Lys Glu Lys Lys Ser Leu Thr Trp Glu Thr Ser Ser Tyr Asp 460 465 455 Leu Glu; and Met Ser Lys Val Lys Leu Thr Lys Glu Asn Ile Val Ser Leu Leu 5 Thr Gln Ser Ala Asp Val Glu Phe Glu Glu Asp Gln Asn Gln Val 25 20 Ala Phe Asn Phe Lys Thr Phe Cys Gln Glu Asn Leu Asp Leu Ile 45 10 35 Lys Lys Met Ser Ile Thr Ser Cys Leu Thr Phe Leu Lys Asn Arg Gln Gly Ile Met Lys Val Val Asn Gln Ser Asp Phe Thr Phe Gly 70 65 Lys Val Thr Ile Lys Lys Asn Ser Glu Arg Val Gly Ala Lys Asp 15 90 85 80 Met Thr Phe Arg Arg Leu Asp Ser Met Ile Arg Val Lys Leu Ile 100 Glu Glu Thr Ala Asn Asn Glu Asn Leu Ala Ile Ile Lys Ala Lys 20 115 120 110 Ile Ala Ser His Pro Leu Val Gln Ala Tyr Gly Leu Pro Leu Ala 135 130 125 Asp Ala Lys Ser Val Arg Leu Ala Ile Met Leu Gly Gly Ser Ile 150 Pro Leu Ile Ala Ser Val Asp Ser Phe Glu Met Ile Ser Val Val 25 165 160 155 Leu Ala Ile Tyr Gln Asp Ala Lys Tyr Lys Glu Leu Gly Ile Glu 175 180 170 Pro Thr Lys Tyr Asn Thr Lys Glu Ala Leu Gly Lys Val Cys Thr 30 Val Leu Lys Ser Lys Gly Phe Thr Met Asp Asp Ala Gln Ile Asn 210 200 205 Lys Gly Lys Glu Tyr Ala Lys Ile Leu Ser Ser Cys Asn Pro Asn 225 220 215 Ala Lys Gly Ser Ile Ala Met Asp Tyr Tyr Ser Asp Asn Leu Asp 35 235 Lys Phe Tyr Glu Met Phe Gly Val Lys Lys Glu Ala Lys Ile Ala 255 245 250 Gly Val Ala As the nucleocapsid protein gene depicted above is on the viral 40 complementary strand, the nucleocapsid protein gene of TSWV-B is: 45 ATG TOT AAG GTC AAG CTC ACA AAA GAA AAC ATT GTC TOT CTT TTA ACT CAA TCT GCA GAT GTT GAG TIT GAA GAA GAC CAG AAC CAG GTC 90

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GCA TIC AAC TIT AAG ACT TIC TGT CAG GAA AAT CIT GAC CIG ATT 135 AAG AAA ATG AGT ATC ACT TCA TGT TTG ACT TTC TTG AAG AAT CGC 180 CAA GGC ATT ATG AAA GIT GIG AAC CAA AGT GAT TIT ACT TIT GGT 225 AAG GIC ACG ATA AAG AAA AAT TCT GAA AGA GIT GGA GCT AAG GAT 270 ATG ACT TIC AGG AGG CIT GAT AGC ATG ATA AGA GIG AAG CIC ATA 315 GAA GAG ACT GCA AAC AAT GAG AAT CIT GCT ATT ATC AAA GCA AAA 360 ATT GOO TOO CAC OOT TIG GIC CAA GOT TAC GGG CIG COT CIG GOA 405 GAT GCA AAA TCT GIG AGA CIT GCT ATA ATG CTT GGA GGT AGT ATC 450 CCT CTC ATT GCT TCT GTT GAC AGC TTC GAG ATG ATC AGT GTT GTT 495 CTT GCC ATA TAT CAA GAT GCA AAG TAC AAG GAG TTA GGG ATT GAA 540 CCA ACT AAG TAC AAC ACT AAG GAA GCT CTG GGG AAG GTT TGC ACA 585 GTG CTT AAA AGC AAA GGA TTT ACA ATG GAT GAT GCA CAG ATA AAC 630 AAA GGG AAA GAA TAT GCC AAG ATA CTT AGT TCT TGC AAT CCC AAT 675 GCT AAG GGA AGC ATT GCT ATG GAC TAT TAC AGT GAT AAT CIT GAC 720 AAA TIC TAT GAA AIG TIT GGA GIT AAG AAA GAG GCC AAG ATT GCT 765 GGT GTT GCA TAA 777

The compete S RNA of TSWV-B should be 3049 nucleotides in length, 134 nucleotides longer than S RNA of TSWV-CPNH1. This difference was mainly attributed to the elongated intergenic region of the TSWV-B S RNA. Analysis of the sequenced region of TSWV-B S RNA revealed two open reading frames as depicted above, which is similar to other TSWV isolates. The larger one was localized on the viral RNA strand originating at nucleotide 88 and terminating at nucleotide 1491. The smaller one on the vial complementary strand was defined by an initiation codon at nucleotide 2898 and a termination codon at nucleotide 2122. The two open reading frames were separated by an intergenic region of 629 nucleotides. Comparisons of the entire sequenced TSWV-B S RNA with S RNA regions of other isolates in the following table which depicts the percent homology comparison of aligned nucleotide and amino acid sequences of the TSWV-B S RNA with those of the other isolates:

		Overall	53 K	protein gene	Intergenic	29 K	protein
	gene						
	Comparisons ^a	n t	n t	aa	n t	n t	a a
	B/CPNH1	76.4 ^b	80.0	86.1(78.3) ^c	72.4	77.5	91.5(79.1)
5	B/L3	75.8	79.0	89.0(82.0)	76.4	78.0	91.1(79.9)
	B/BL	76.3	-	•	72.8	77.6	90.3(79.5)
	B/I	63.0	-	-	-	63.1	69.7(55.3)
	CPNH1/L3	94.8	95.6	92.0(89.4)	89.2	96.8	99.6(98.5)
	CPNH1/BL	96.4	-	-	95.9	97.2	98.8(96.9)
10	CPNH1/I	62.7	-	-	•	60.8	69.5(55.1)
	L3/BL	95.1	-	-	92.6	97.3	99.2(98.5)
	L3/I	60.9	-	-	•	60.9	69.5(55.1)
	I/BL	61.7	-	-	-	60.9	68.8(53.9)

The partial or complete S RNA sequences of isolates TSWV-CPNH1 (2.916 kb), TSWV-L3 (2.837 kb), TSWV-BL (2.037 kb) and TSWV-I (1.144 kb) were used for comparisons with the S RNA sequence of the TSWV-B (3.049 kb).

20 c Percent identity is in parenthesis.

As depicted, the greatest nucleotide sequence similarity (75.8%-76.4%) was shown with the L-type isolates (CHNH1, L3 and BL). To the lesser extent, there was nucleotide sequence similarity (63%) between the TSWV-B S RNA and the S RNA of TSWV-I assigned to I serogroup.

For comparison, the sequenced S RNA regions of the L-type isolates (CHPN1, L3 and BL) shared 94.8%-96.4% nucleotide sequence similarities.

The open reading frame of 777 nucleotides encodes the N protein of 258 amino acids with a predicted molecular weight of 28700 Da. the sequence comparisons of the N open reading frame from TSWV isolates revealed that nucleotide sequences of the N genes from the isolates CPNH1, L3 and BL differs from TSWV-B by a considerably larger amount (22%-22.5%) than they differ from each other (2.7%-3.2%). Consistent to the results of the immunological analysis, the N amino acid

sequences among CPNH1, L3 and BL isolates are more closely related to each other (98.8%-99.6% similarities or 96.9%-98.5% identities) than to

b Percent similarities were calculated by Comparison of their nucleotide or predicted amino acid sequence using the program BESTFIT of the GCG Sequence analysis software package.

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the TSWV-B (90.3%-91.5% similarities or 79.1%-79.9% identities). Much lower homology was observed to TSWV-I at both nucleotide (63.1%) and amino acid (69.7% similarity or 55.3% identity) levels. Except for the N open reading frame of TSWV-I that encodes 262 amino acids, the N open reading frames of the other isolates code for the 258 amino acids. Computer analysis suggested that the extra residues of TSWV-I N open reading frame resulted from the amino acid sequence insertions (residues 82 through 84 and residue 116). One potential N-glycosylation site is found at residue 68.

The second open reading frame of 1404 nucleotides encodes the nonstructural protein of 467 amino acids with a predicted molecular weight of 52566 Da. Comparisons with homologous open reading frames of TSWV-CPNH1 and TSWV-L3 showed 80% and 79% similarities at the nucleotide level, and 86.1% (or 78.3% identity) and 89% (or 82.0% identity) similarities at the amino acid level. This open reading frame contains four potential glycosylation sites, which are located in the exactly same positions as those of TSWV-CPNH1 and TSWV-L3.

The intergenic region of the TSWV-B S RNA was, due to several insertions, 126 and 41 nucleotide longer than the counterparts of TSWV-CPNH1 and TSWV-L3, respectively. The sequence analysis by the program FOLD indicated the intergenic region can form very complex and stable hairpin structure by internally base-pairing U-rich stretches with A-rich stretches of the intergenic region, which had similar stability to those produced from TSWV-CPNH1 and TSWV-L3 as indicated by minimum free energy values. This internal base-paired structure may act as a transcription termination signal.

The results tabulated above also revealed that the N protein of TSWV-B is subject to a higher degree of selective pressure than the 52 K protein; the similarities among the amino acid sequences of the 52 K protein are lower than that found for the amino acid sequence of the NPs. Nucleotide sequence divergence is highest among the intergenic regions, which indicates that this region is subject to less selective pressure than either genetic region.

The evolutionary relationships among the TSWV-B and other four TSWV isolates were analyzed and depicted in figure 4 in which the

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evolutionary tree organization is consistent with the relatedness of serological data collected for these TSWV isolates. Thus, the TSWV-B, according to the present invention, is more closely related to the L-type isolates than to the I-type isolate TSWV-I, but is much less similar to the L-type isolates than the L-type isolates are to each other.

Despite a slight delay of symptom expression, transgenic plants did not show resistance to the Brazil isolate of TSWV; Serological results show that this isolate is distinct from the "L" and "I" type isolates, and biologically different from the curcurbit isolate. The Brazil isolate may thus belong to still another serogroup of TSWV. In any event, infectivity results show that it is unlikely that a single NP gene will provide resistance to all isolates in the Tospovirus genus.

Transgenic plants according to the present invention that gave low or undetectable ELISA reactions (0-0.05 OD405nm) were resistant to infection by the heterologous isolates (Arkansas and 10W pakchoy) of the "L" serogroup, whereas no protection against these isolates was found in plants accumulating high levels of the NP. Compared to the ELISA readings of control NP(-) plants (0.05 OD405nm), these transgenic plants may produce little, if any, TSWV-BL NP. Similar

results have been observed in transgenic plants, in which the CP accumulation was not detected; these were highly resistant to virus infection. The mechanism underlying this phenomenon is presently unknown. It is likely that this type of resistance might be attributed to interference of CP RNA molecules produced in transgenic plants with viral replication, presumably by hybridizing to minus-sense replicating RNA of the attacking virus, binding to essential host factors (e.g., replicase) or interfering with virion assembly.

It should be noted, however, that the resistance to the homologous TSWV-BL isolate is apparently independent of the expression levels of the NP gene. Although the relative NP levels of the individual R₁ plants inoculated with TSWV-BL were not measured, it is reasonable to assume that the NP produced in these inoculated R₁ plants (a total of 145 plants tested) ranged from undetectable to high.

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In contrast to the case for protection against the heterologous isolates of the "L" serogroup, protection against the Begonia isolate of the TSWV-I serogroup was found in the high NP-expressing R1 plants. Comparison of NP nucleotide sequence of the "L" serogroup with that of the "I" serogroup revealed 62% and 67% identity at the nucleotide and amino acid levels, respectively. The difference of NP genes of the two serogroups might be so great that the NP (the "L" serogroup) produced in transgenic plants acted as a dysfunctional protein on the attacking Begonia isolate of the "I" serogroup. Incorporation of this "defective" coat protein into virions may generated defective virus which inhibit virus movement or its further replication. This type of interaction is expected to require high levels of the NP for the protection. Alternatively, resistance to the Begonia isolate may also involve interference of NP transcripts produced in R₁ plants with viral replication. If this is true, more NP transcripts (due to the heterologous nature of two NP gene) may be required to inhibit replication of heterologous virus.

Although there are no obvious explanations for the results showing the relation of NP levels in individual R₁ plants to resistance to the heterologous isolates of the "L" and "I" serogroups, it is believed these are definite trends since the data were derived from a large number (190) of plants. Thus, it is believed that a measurement of CP or NP levels in individual plants may provide a more accurate way to relate NP or CP levels to resistance. By this form of data analysis, the results show that the resistance was more closely related to NP levels in each test plant than to the NP level of the R₀ line from which they were derived. For TSWV-BL Np gene in tobacco, at least, it appears that integration sites of the NP gene in plant chromosome may not be important for viral resistance.

Studies have also been conducted to determine the reaction of transgenic R₁ and R₂ tomatoes containing the nucleocapsid protein gene of TSWV-BL according to the present invention to the following isolates: Brazil (a distantly related virus), T91 (a closely related virus) and BL (a homologous isolate). In these studies, transgenic tomatoes (*L. esculentum*) were produced by *A. tumefaciens*-mediated gene transfer

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of the nucleocapsid protein (N) gene of the lettuce isolate of tomato spotted wilt virus BL into germinated cotyledons using modifications of published procedures [see Plant Cell Reports 5:81 (1986)]. The tomato line "Geneva 80" was selected for transformation because it contains the Tm-22 gene which imparts resistance to TMV, thus creating the possibility of producing a multiple virus-resistant line.

Transformants were selected on kanamycin media and rooted transgenic tomatoes were potted and transferred into the greenhouse. R₁ and R₂ tomato seedlings expressed the NPT II gene, suggesting multiple insertions of this gene in the plant genome. In contrast, only 18% of the seedlings produced detectable levels of the N protein.

Nine R₁ and three R₂ lines were tested for resistance to the following three *Tospovirus* described, specifically TSWV-BL, TSWV-T91, and TSWV-B. Infectivity was based upon visual inspection of test plants. In those cases where plants appeared healthy except for a few rust-colored rings or insect damage, extracts from these plants were inoculated to N. benthamiana to test for the presence of the virus. As depicted in the following table, nearly all control tomato plants exhibited typical symptoms consisting of plant stunting, leaf yellow mosaic and rugosity 3 to 4 weeks after inoculations with TSWV-BL, TSWV-T91 or TSWV-B. However, only 4% of the R₁ and R₂ transgenic plants became infected with TSWV-BL, 7% with TSWV-T91, and 45% with TSWV-B.

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Viral resistance in transgenic R1 and R2 tomatoes expressing the nucleoprotein gene of the lettuce strain of tomato spotted wilt virus

		Inocu	lating Isolate	_{es} a
	Plant Line	<u>ŤSŴV-BL</u>	TSWV-T91	TSWV-B
5	R1 Plants:			
	T13-1	0/22	1/26	7/24
	†13-2	6/20	ИТР	NT
	T13-3	2/42	0/20	12/18
	T13-4	0/25	NT	NT
10	T13-9	0/20	NT	NT
	T13-10	1/50	2/26	11/26
	T13-11	0/22	NT	NT
	T13-12	1/29	NT	NT
	T13-13	0/22	NT	NT
15	TOTAL	10/252	3/72	30/68
	R2 Plants:			
	T13-1-7	0/8	2/8	5/8
	T13-1-9	0/8	1/8	2/8
	T13-1-11	0/8	1/9	5/9
20	TOTAL	0/24	4/25	12/25
	CONTROLS	92/95	51/53	52/53

plants were inoculated at the one- to two-leaf stage with 5-, 10-, or 20fold diluted leaf extract of N. benthamiana, H423 tobacco or tomato; the same plants were re-inoculated 7 days later and symptoms were recorded after another 14 days; the reaction is expressed as number of plants with symptoms/number of plants tested

b not tested

Accordingly, the description above supports the finding that transgenic tomato plants that express the N gene of TSWV-BL show resistance to infection to TSWV-BL, to other TSWV isolates that are closely related to TSWV-BL, and to the more distantly related TSWV-B.

In further limited studies with an additional isolate, all transgenic plants were resistant to the 10W (pakchoy) isolate, whereas the controls were infected. These results show that transgenic tomatoes are better protected against closely related isolates than distantly related isolates. Unlike in transgenic tobacco and N. benthamiana expressing the TSWV-BL N gene, the level of N protein expression did not correlate with the observed protection in transgenic

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tomatoes; 55% of the transgenic tomatoes were also resistant to a distantly related isolate of TSWV-B, which was not observed in transgenic tobacco and N. benthamiana plants. These discrepancies may reflect that tomato is inherently less susceptible to Tospoviruses.

In addition, studies were also conducted to determine virus distribution in a small number of plants at 5 and 7 weeks after inoculation. The distal halves from leaflets of all expanded leaves of each plant were ground and back-inoculated onto N. benthamiana. The results taken seven days after inoculation showed that virus cannot be recovered from any leaf tissue of asymptomatic transgenic plants inoculated with either TSWV-BL. -T91, or -B, confirming the visual findings reported above. In transgenic plants showing symptoms, the virus is not distributed throughout the plant. For example, a transgenic plant which could not be conclusively rated visually contained the virus in only two of the 8 leaves; the second leaves from the bottom and top of the plant. Conversely, virus present in all leaves of the infected control plant, and is absent in those of the healthy control plants.

Graft inoculations were attempted to test whether the resistant transgenic plants could become infected if virus is introduced into the vascular system. R1 and R2 plants that had been inoculated at 1:5, 1:10 or 1:20 dilutions of TSWV-BL, -T91, or -B were grafted onto control plants infected with the same isolates and dilutions. The 34 transgenic plants were asymptomatic after 31 days, although the non-transgenic controls were infected. After 23 days, the top 46 cm of transgenic plants had been trimmed away to induce new growth and more plant stress. Although the young, vigorously growing new shoots failed to show any symptoms on the 31st day post inoculation, 33%, 31% and 45% of TSWV-BL, -T91 and -B were showing leaf or stem symptoms, respectively at 45 days post inoculation. These results indicate that some transgenic plants are tolerant, and others are immune to infection.

Thus, according to one aspect of the present invention, transgenic plants expressing the NP gene of the TSWV-BL isolate are highly resistant to infections of both the homologous TSWV-BL isolate and heterologous isolates of the same serogroup (Arkansas and 10W

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pakchoy). More significantly, the resistance is effective to Begonia isolate from other serogroups. In brief, the above clearly describes that transgenic tobacco plants expressing the nucleoprotein gene of TSWV-BL display resistance to both TSWV and INSV, and the protection appears to be mediated by the nucleoprotein against distantly related INSV and by the nucleoprotein gene ribonucleotide sequence against the homologous and closely related TSWV isolates. This is the first time broad spectrum resistance of the engineered plants to different isolates of TSWV has been shown.

While coat protein protection generally displays delay and/or reduction in infection and symptom expression, but no immunity, the present invention provided a significantly high percentage of transgenic plants which were symptom-free and free of the infective virus. Resistance of these plants under greenhouse conditions persisted throughout their life cycle, and more importantly was inherited to their progenies as shown above.

It was observed in the present invention that the transgenic plants producing little, if any, TWSV-BL NP were highly resistant to infection by the homologous isolate and other closely-related isolates within the same serogroup of TSWV, whereas no protection was found in those expressing high levels of the NP gene.

The biological diversity of TSWV is well documented and has been reported to overcome the genetic resistance in cultivated plants such as tomato. Thus, it is extremely important to develop transgenic plants that show resistant to many strains of TSWV. The present invention indicates that one method to do so would be to utilize the viral NP gene to confer this resistance, and that this resistance would be to diverse TSWV isolates. Thus, the finding of the present invention that the expression of TSWV NP gene is capable of conferring high levels of resistance to various TSWV isolates has a great deal of commercial importance.

In another series of studies, Plasmid BIN19-N+ was constructed and transferred to A. tumefaciens strain LBA4404 in accordance with Example IV, and transferred to Nicotiana benthamiana in accordance

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with Example V. The nucleocapsid genes of INSV-Beg and -LI were amplified with oligomer primers INSV-A (5'-TAGTTATOTAGAACCATGGACAAAGCAAAGATTACCAAGG) and INSV-B (5'-TAGAGTGGATCGATGGTTATTTCAAATAATTTATAAAAGCAC),

5 hybridizing to the 5'-coding and 3'-noncoding regions of the nucleocapsid gene of an INSV isolate, respectively. The amplified nucleocapsid gene fragments were purified in accordance with Example III, and digested and sequenced in accordance with Example IV.

Of a total of 24 N+ (transformed with pBIN19-N+) and 18 N-(transformed with vector pBIN19) transgenic N. benthamiana plants were transferred to soil and grown in the greenhouse. All N+ lines were confirmed by PCR at leaf stages 4-5 to contain the N gene sequence. The relative level of N protein accumulation was estimated in each independent Ro transgenic clonal line by DAS-ELISA using antibodies of the TSWV-BL N protein. Of the twenty-four N+ lines, two had OD405nm readings of 0.50-1.00, seventeen between 0.02-0.10, and the remaining five less than 0.02. Healthy N. benthamiana or transgenic N- plants gave OD405nm readings of 0.00-0/02. All the Ro plants were selfpollinated and the seeds from the following transgenic lines were germinated on kanamycin (300 µg/ml) selection medium for inoculation tests: (1) N- -2 and -6, control transgenic lines containing vector pBIN19 alone; (2) N+-28, a transgenic line that produced an undetectable amount of the N protein (OD405nm = 0.005); (3) N+-21, a transgenic line producing a low level of the N protein (OD405nm = 0.085); and (4) N+-34 and -37, two transgenic lines accumulating high

0.085); and (4) N+-34 and -37, two transgenic lines accumulating high levels of the N protein (OD405nm = 0.50-1.00. These six lines were then analyzed by Northern hybridization; the intensity of N gene transcripts correlated well with the levels of ELISA reactions.

Transgenic seedlings from the six R₀ lines were selected by
germinating seeds on kanamycin selection medium, and these seedlings
were inoculated with the five *Tospoviruses*. The inoculated R₁ plants
were fated susceptible if virus symptoms were observed on
uninoculated leaves. In order to exclude the possibilities of escapes,
transgenic control N- plants were always used in each inoculation of
transgenic N+ plants. In addition, each inoculum extract was always

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used to first inoculate N⁺ plants followed by control N⁻ plants. The results from this series of studies are depicted below:

Reactions of R1 plants expressing the nucleocapsid (N) protein gene of N. benthamiana spotted will virus (TSWV) to inoculation with Tospoviruses

Š No. plants infécted/No. plants inoculated

			TSWV	SOLATE	INSV ISC	DLATE	
	Ro Line	ELISAa	B L	10W	Beg	LI	TSWV-B
	N2/-6	<0.02	32/32	32/32	32/32	20/20	32/32
	N+-28	0.005	16/16	16/16	15/16		16/16
lO	N+-21	0.085	9/40	17/40	39/40	, 18/20	40/40
,	N+-34	0.715	25/28 ^C	28/28	23/28 ^C	·	28/28
	N+-37	0.510	26/28 ^c	22/22	21/28 ⁰	16/20 ^C	22/22

aELISA data of Ro lines from which the Ro plants were derived;

b30-föld diluted léaf extracts of infected N. benthamiana plants were applied to the three leaves of plants at the 3-5 leaf stages. Each extract was always used to inoculate N+ plants followed by control N- plants. Data were taken daily for at least two months after inoculation and expressed as number of plants systemically infected/number of plants inoculated;

cindicate that nearly all susceptible R1 plants displayed a significant delay of symptom appearance.

As depicted in the above table, all R1 plants from control lines N-2 and -6 showed systemic symptoms 5-8 days after inoculation with all the viruses tested. None of the R1 plants from line N+-28 produced detectable levels of the N protein, and all were susceptible to these viruses except for one plant inoculated with INSV-Beg. ELISA assays of leaf discs from this N+-28 R1 plant sampled before inoculation clearly showed that the plant identified to possess the INSV-Beg resistant phenotype did accumulate a high level of the N protein (OD405nm = 0.78 as compared to OD405nm <0.02 for all other N+-28 R1 plants).

The low N gene expressing line N+-21 showed the best resistance against the homologous (78%) and closely related TSWV-10W (57%) isolates and very little resistance to the two INSV isolates (3% and 10%); only three N+-21 plants showed the resistant phenotype when inoculated with the INSV isolates. Leaf samples from these INSV-resistant N+-21 R₁ plants gave much higher ELISA reactions (OD405nm 0.5 to 1.00) and thus higher amounts of the N protein than the

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susceptible N+-21 plants (OD405nm 0.02 to 0.20). The high N gene expressing lines N+-34 and -37 showed the highest resistance to INSV isolates (18%-25%) followed by the homologous TSWV-BL isolate (7% and 11%) while none of the plants showed resistance to TSWV-10W; however, the N+-34 and -37 R1 plants that became infected with INSV or TSWV-BL did show various lengths of delays in symptom expression. None of the R1 plants from these four transgenic N+ lines were resistant to TSWV-B; some of the R1 plants from the N+-34 and -3,7 lines showed a slight delay of symptom appearance

In studies to determine whether the level of N protein production in N+ R1 plants was associated with resistance to different Tospoviruses, the inoculated N+ R1 plants in the preceding table were te-organized into four groups based on the intensity of their ELISA reactions of tissues taken before inoculation irrespective of original Ro pants. The N+ R1 plants that expressed low levels of the N protein (0.02-0.2 OD) showed high resistance (100% and 80%) to TSWV-BL and -10W but were all susceptible to INSV-Beg and -LI, showing no detectable delay in symptom expression relative to control N- plants. In contrast, nearly all N+ R1 plants with high levels of the N protein (0.20-1.00 OD) showed various levels of protection against TSWV-BL, INSV-Beg and -LI, ranging from a short delay of symptom expression to complete resistance with most of these plants showing various lengths of delay in symptom development relative to control N- plants. No protection was observed in the high expressors against TSWV-10W. In addition, none of the N+ R₁ plants were resistant to TSWV-B regardless of the level of N gene expression; however, a short delayed symptom appearance was observed in the N+ R1 plants producing high levels of the N protein. All control N-R1 plants and transgenic N+R1 plants with undetectable ELISA reactions (0 to 0.02 OD) were susceptible to all the Tospoviruses tested.

The inhibition of replication of a distantly related INSV in N. benthamiana protoplasts expressing the TSWV-BL nucleocapsid gene was also studied. In these studies, whole INSV-LI virions were used to infect protoplasts that were isolated from three transgenic lines to investigate how the products of the transgene affect replication of the

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incoming virus. Viral replication was determined by measuring the lével of the N protein of the infecting INSV in transgenic protoplasts using antibodies specific to the INSV N protein. DAS-ELISA analysis showed that all progenies from a given line were relatively uniform and nearly all R1 progeny gave an expression level of transgenic N gene similar to their parental transgenic line. These results allowed for the prediction of the expression level of R1 populations based on that of their parental lines. Protoplasts derived from R1 plants of the low expressor line N+-21 supported the replication of INSV-LI whereas protoplasts from R1 plants of the higher expressor line N+-37 did not until 42 hours after inoculation at which low levels of viral replication Were observed. The same protoplasts at various time intervals (e.g. 0, 19, 30 and 42 hours) were also assayed by DAS-ELISA using antibodies specific to the TSWV-BL N protein to monitor the expression level of the transgene. As expected, protoplast from N+-21 R1 plants produced relatively low levels (0.338-0.395 OD405nm) whereas protoplasts from N+-37 R₁ plants accumulated high levels (0.822-0.865 OD405nm). The expression level was found to be consistent at all time points.

In this aspect of the present invention it has been shown that transgenic N. benthamiana plants that accumulate low amounts of the TSWV-BL N protein are highly resistant to the homologous and closely related (TSWV-10W) Isolates, while plants that accumulate high amounts of this protein posses moderate levels of protection against both the homologous and distantly related (INSV-Beg and INSV-LI) viruses. More importantly, these findings indicate that transgenic N.

benthamiana plants (a systemic host of INSV) are protected against INSV-Beg and INSV-LI isolates.

As discussed above, we have shown that transgenic plants expressing the N gene of TSWV are resistant to homologous isolates, and that such plants expressing the TSWV-BL N gene are resistant to both TSWV and INSV. It has also been shown the best resistance to homologous and closely related isolates was found in transgenic plants accumulating low levels of N protein while transgenic plants with high levels of TSWV-BL N protein were more resistant to serologically

35 distant INSV isolates. This observation led us to suspect the role of

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the translated N protein product in the observed protection against homologous and closely related isolates and to speculate that either the N gene itself which was inserted into the plant genome or its transcript was involved in the protection. To test this hypothesis transgenic plants containing the promoterless N gene or expressing the sense or antisense untranslatable N coding sequence were produced. What was discovered was that both sense and antisense untranslatable N gene RNAs provided protection against homologous and closely related, isolates, and that these RNA-mediated protections were most effective in plants that synthesized low levels of the respective RNA species and appears to be achieved through the inhibition of viral replication.

More specifically, the coding sequences introduced into transgenic plants is shown in figure 7. As depicted, the construct pBIN19-N contains the promoteriess N gene inserted into the plant transformation vector pBIN19 (see Example IV). All other constructs contain a double 35S promoter of CaMV, a 5'-untranslated leader sequence of alfalfa mosaic virus and a 3'-untranslated/polyadenylation sequence of the nopaline synthase gene. pBI525 is a plant expression vector and is used in this study as a control; pBI525-mN contains the mutant (untranslatable) form of the N gene; pBI525-asN contains the antisense form of the untranslatable N gene. One nucleotide deletion at the 5'-terminus of the mutant N gene is indicated by the dash symbol. ATG codons are underlined and inframe termination codons in the mutant gene are shown in bold.

EXAMPLE VIII

Primer-directed mutagenesis and cloning of the TSWV-BL N gene was conducted as follows:

Full-length N gene was obtained by reverse transcription and polymerase chain reaction as described in Phytopathology 82:1223 (1992), the disclosure of which is incorporated in toto herein. The untranslatable N coding sequence was similarly generated by RT-PCR using oligomer primers A (AGCATTGGATCATGGTTAACACACTAAGCAAGCAC), which is identical to the S RNA in the 3'-noncoding region of the TSWV-BL N gene, and B (AGCTAATCTAGAACCATGGATGACTCACTAAGGAAAGCATTGTTGC).

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complementary to the S RNA in the 5'-terminus of the N gene. The latter oligomer primer contains a frameshift mutation immediately after the translation initiation codon and several termination codons to · block possible translation readthroughs. The intact and mutant N gene fragments were purified on a 1.2% agarose gel as described in Example II. The gel-isolated intact and mutant N gene fragments were digested with the appropriate restriction enzyme(s) and directly cloned into BamHI/Xbal-digested plant transformation vector pBIN19 and Ncoldigested plant expression vector pBI525, respectively as described in Example IV. The resulting plasmids were identified and designated as pBIN19-N containing the intact, promoterless N gene, and pBI525-mN and pBI525-asN containing the mutant coding sequence in the sense and antisense orientations, respectively, relative to cauliflower mosaic virus 355 promoter. The translatability of the mutant N coding sequence in the expression cassette was checked by transient expression assay in Nicotiana tabacum protoplasts; and the expression cassettes containing the sense or antisense mutant N coding sequence were then excised from plasmid pBI525 by a partial digestion with HindIII/Ecori (since the N coding sequence contains internal HindIII and ExoRI sites), and ligated into the plant transformation vector pBIN19 that had been cut with the same enzymes. The resulting vectors as well as pBIN19-N were transferred to A. tumefaciens strain LBA4404 using the procedure described in Example IV. Leaf discs of N. tabacum var Havana cv 423 were inoculated with the A. tumefaciens strain LBA4404 containing various constructs and the resulting transgenic plants were self-pollinated and seeds were selectively germinated on kanamycin medium.

PCR was performed on each R₀ transgenic line as described above. The oligomer primers A and B were used to determine the presence of the N coding sequence of TSWV-BL. The oligomer primer 35S-promoter (CCCACTATCCTTCGCAAGACCC) was combined with either the oligomer primer A or B to confirm the orientation (relative to the CaMV 35S promoter) of the mutant N coding sequence inserted into the plant genome. DAS-ELISA used to detect the N protein in transgenic plants was performed using polyclonal antibodies against the TSWV-BL

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N protein. For an estimation of RNA transcript level in transgenic plants by Northern blot, total plant RNAs were isolated according to Napoli [see The Plant Gell 2:279 (1990)], and were separated on a formaldehyde-containing agarose gel (10 µg/lane). The agarose gels were then stained with ethidium bromide to ensure uniformality of total plant RNAs in each lane. Hybridization conditions were as described in the GeneScreen Plus protocol by the manufacturer. Resulting signal blots were compared and normalized based on the N gene transcript band of the control lane (the mN R₁ plant producing a high level of the N gene transcript) included in each blot. The transgenic plants that gave density readings (Hewlet ScanJet and Image Analysis Program) between 100 and 150 were rated as high expressors, while the plants with densities between 15 and 50 were rated as low expressors.

Inoculation of transgenic plants with *Tospovirus* was carried out as described above with inoculation being done at the 3-4 leaf stage except were indicated.

Tobacco protoplasts were prepared from surface-sterilized leaves derived from R1 plants [see Z. Pflanzanphysiol. 78:453 (1992) with modifications]. The isolated protoplasts (6 x 106 protoplasts) were transformed with 0.68 OD260nm of the purified TSWV-BL virion preparation using the PEG method [see Plant Mol. Biol. 8:363 (1987)]. The transformed protoplasts were then cultured at the final density of 1 x 106 protoplasts /ml in the culture medium at 26°C in the dark.

After various intervals of incubation, the cultured protoplasts were washed twice with W5 solution and lysed by osmotic shock in the enzyme conjugate buffer. Viral multiplication (replication) was estimated by measuring the N protein of the virus using DAS-ELISA.

As described, one aspect of the present invention demonstrated that transgenic tobacco producing none or barely detectable amounts of the N protein were resistant to homologous and closely related isolates. This result suggested that the observed resistance may have been due to trans interactions of the incoming viral N gene RNA with either the N gene transcript produced in the transgenic plants or the N coding sequence itself. To test whether the presence of the nuclear N gene

plays a role, transgenic PoN Ro lines and Ro plants from two PoN lines were challenged with four *Tospoviruses* (TSWV-BL, TSWV-10W, INSV-Beg and TSWV-B). Only asymptomatic plants were rated resistant while plants showing any symptoms were rated susceptible. All inoculated Ro and Ro plants were susceptible to the viruses.

To further test the possibility that the transcript of the N transgene is involved in the protection, a number of R₀ transgenic plants that produced either the sense or the antisense N gene transcript but not the N protein were inoculated with the homologous isolate.

10 Results appear in the following table:

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	Form of transgenea	Level of N gene RNAb	No. of R0 lines tested	No. of lines inoculated ^c	No. of lines resistant
	mΝ	Н	8	4	0
		L	17	16	16
5		nd	4	1	0
	asN	H	6	3	0
		L	9	5	5
		nd	1	0	0
	P°N	nd	12	6	0

- amN and asN represent plants expressing the sense and antisense untranslatable N genes, respectively, P°N represents plants containing the promoterless N gene; bithe level of the N gene RNA was estimated in each line by Northern blots, nd indicates that the N gene transcript was not detected;
- 25 C30-fold diluted leaf extracts of the N. benthamiana plants infected with TSWV-BL were applied to three leaves of each plant at the 6-7 leaf stage. Each extract was first applied to all test plants followed by control healthy plants. Data were taken daily for 45 days after inoculation and only the asymptomatic plants were rated resistant.
- Unlike the controls, which developed typical systemic symptoms 7 to 9 days after inoculation, 16 out of 21 mN plants and 5 our of 8 asN plants were asymptomatic throughout their life cycles. Northern blot analysis of leaf tissues sampled before inoculation showed that all the resistant R₀ lines produced low levels of the sense or antisense N gene RNA, whereas the susceptible R₀ lines produced either none or high
- levels of the RNA species. Since this data suggested that the resistance of transgenic plants to TSWV-BL was related to their relative levels of N gene transcript, transgenic progenies from four mN

and three asN Ro lines with either high or low N gene transcript levels were selected by germination on kanamycin-containing media. These transgenic plants were tested for resistance to the four *Tospoviruses* at the 3 to 4 leaf stage, except that some R₁ plants from two asN lines were inoculated at the 6 to 7 leaf stage. The results are summarized in the following table:

Ro Line	N gene RNAa	TSWV-BL	TSWV-10W	INSV-Beg	TSWV-B
Promoterless N gene	ine				
P°N-1	1 2	10/10	10/10	10/10	10/10
P°N-2	þ	15/15	10/10	10/10	10/10
N°-3	2	8/8	9/9	9/9	9/9
Untranslatable N gene	lene				
mN-2	I	20/20	20/20	20/20	20/20
1-Nm	I	20/20	20/20	20/20	20/20
BN-13	_1	2/20	4/20	20/20	20/20
mN-18	٦	4/20	1/20	20/20	20/20
N°-3	.	24/24	32/32	24/24	24/24
Antisense N gene					
asN-1	ب	20/20 ^b	20/20	20/20	20/20
asN-4	I	20/20	20/20	20/20	20/20
		(16/16) ^C	(16/16)		
asN-9	ب	19/20 (3/41)	<u>20/20</u> (5/21)	20/20	20/20
N°-3	1 2	16/16 (32/32)	16/16 (20/20)	16/16	16/16

aNorthern análysis of Ro lines from which the R1 plants were derived (see preceding table);

bithe underlined fractions indicate that most of susceptible R1 plants displayed a significant delay of symptom appearance;

City fraction in parenthesis represents the inoculation data obtained from plants inoculated at the 6-7 leaf stage; the remaining data in this table were generated from plants inoculated at the 3-4 leaf stage; inoculated plants were observed daily for 45 days after inoculation.

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All R1 plants from high expressor lines mN-2 and mN-7 were susceptible to infections by all Tospoviruses tested, and these plants did not show a delay of symptom appearance as compared to controls. In contrast, high proportions of the R1 plants from low expressor lines mN-13 and -18 were resistant to homologous (TSWV-BL) and closely related (TSWV-10W) Isolates, but not resistant to infections by distantly related Tospoviruses (INSV-Beg and TSWV-B). The resistance of asN R1 plants from low expressor R0 lines was markedly influenced by the TSWV isolate used for inoculation. All but one of the small R1 plants (3-4 leaf stage) from low expressor lines asN-1 and -9 became infected, although there was a delay of symptom appearance, when inoculated with the homologous TSWV-BL or closely related TSWV-10W isolates. In contrast, most of the large R1 plants (6-7 leaf stage) from line asN-9 were resistant to both isolates. In comparison, control R1 plants and R1 plants from the high expressor line such as as N-4 displayed no resistance to either of the isolates regardless of the size of test plants. Antisense RNA-mediated protection was not effective against infection by the distantly related INSV-Beg and TSWV-B isolates.

Analyses of data presented in the above two tables suggest that sense and antisense RNA-mediated protections are observed only in low expressors of the N gene. The R1 asN plants that produced high levels of the antisense N gene transcript were as susceptible as control plants. In contrast, the asN low expressors displayed a delay in symptom appearance when inoculated at the 3-4 leaf stage and showed increased levels of resistant when inoculated at the 6-7 leaf stage.

Inhibition of viral replication in tobacco protoplasts expressing the sense or antisense form of untranslatable N coding sequence was also noted. In this instance, whole virion preparations of TSWV-BL were used to transfect protoplasts isolated from transgenic lines to investigate the effect of sense or antisense N gene transcript on replication of the incoming virus. Viral replication was determined by measuring the level of the N protein of the incoming virus in transfected protoplasts, and it was found that protoplasts derived from plants (mN-7 and asN-4) that produced high levels of the respective

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RNA transcripts supported the replication of the virus, whereas protoplasts from mN low expressor (mN-18) did not. Protoplasts from an asN low expressor (asN-9) supported much lower levels of viral replication.

Accordingly, in this aspect of the present invention we have shown that transgenic plants expressing sense or antisense form of uniffansiatable N gene coding sequence are resistant to homologous (TSWV-BL) and closely related (TSWV-10W), but not to distantly related (INSV-Beg and TSWV-B) Tospoviruses. The following table provides a comparison of resistance to Tospoviruses between transgenic tobacco expressing various forms of the TSWV-BL N gene:

	Homology to	Forn	of the	Trans	<u>gene</u> a
<u>Tospovirus</u> .	TSWV-BL N Geneb	N	<u>mN</u>	<u>asN</u>	P°N
TSWV-BL	100%	R	R	Кc	S
TSWV-10W	99%	R	R	Rc	S
INSV-Beg	60%	Кc	S	S	S
TSWV-B	78%	S	S	S	S

areactions of transgenic tobacco and N. benthamiana plants expressing the intact N gene (N) of TSWV-BL to inoculation with the four Tospoviruses are included for comparisons with inoculation results of transgenic plants containing untranslatable (mN), antisense (asN), and promoteriess (P°N) N coding sequences, R = resistant, S = susceptible;

bithé nucleotide sequences are as reported in Phytopathology 82:1223 (1992) and Phytopathology 83:728 (1993)

clevel of resistance may depend upon the concentration of inoculum.

These results confirm and extend the earlier aspects of the present invention for RNA-mediated protection with TSWV. Furthermore, the protection is observed in plants producing low rather than high levels of the N gene transcript, and although earlier studies reported herein indicate that tobacco plants which produced high levels of the TSWV-BL N protein displayed resistance to INSV-Beg, this additional data indicates that since resistance to INSV-Beg was not observed in transgenic plants expressing the sense or antisense form of the unitranslatable of the N gene thus clearly indicating that protection against INSV-Beg is due to the presence of the N protein and not the N gene transcript. Thus, it appears that two different mechanisms are

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Involved in protection transgenic plants against TSWV and INSV Tospoviruses according to the present invention. One mechanism Involves the N gene transcript (ANA-mediated), and another involves the N protein (protein-medicated). In addition, the results of the protoplast experiments indicate that N dene RNA-mediated protection is achieved through a process that inhibits viral replication, and the data contained in the above tables suggest that protection against the distantly related INSV-Beg isolate is conferred by the N protein of TSWV-BI, and not by the gene transcript.

Finally, further studies were conducted to provide still another aspect of the present invention - that a portion of the Tospovirus nucleopfotein gene provide protection of transgenic plants against infection by the Tospovirus. It has already been demonstrated above that the N gene RNA protects against homologous and closely realated TSWV isolates while the N protein protects against the homologous Isolate and distantly related INSV isolates; that N gene RNE-mediated protection is effective in plants expressing low levels of the N gene whereas N protein-mediated protection requires high levels of N protein accumulation; and that the N gene RNA-mediated protection is achieved through inhibition of viral replication. Based upon this prior data, we next set out to determine whether a portion of the N gene might work against infection by the virus. We found, as discussed below, that transgenic plants expressing about one-half of the N gene sequence is resistant to the virus.

The following describes the cloning of one-half N gene fragments of TSWV-BL in order to demonstrate this final aspect of the present invention. The first and second halves of both the translatable and untranslatable N gene were generated by reverse transcription and then PCR as described above. As depicted in figure 8, the nucleotide deletion or insertions at the 5'-terminals of the untranslatable half N dene fragments are indicated by the dash symbol; ATG codons are underlined and all possible termination codons immediately after the initiation codon of the untranslatable half N gene fragments are shown in bold.

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The first half of the N gene was produced by RT-PCR using oligoprimers I (5'-TAGAGTGATCCATGGTTAAGGTAATCCATAGGCTTGAC), which is complementary to the central region of the TSWV-BL N gene, and II (5'-AGCTAAGGATGATTAAGCTCACTAAGGAAAGCATTGTTGC) for the translatable or iii

- (5'-AGGTAATOTAGAAGGATGGATGACTCACTAAGGAAAGCATTGTTGC) for the untranslatable first half N gene fragment, the latter two oligomer primers are identical to the 5'-terminus of the N gene. Similarly, the second half of the N gene was produced by RT-PCR using oligomer
- primers iv (5'-AGCATTGGATCCATGGTTAACACACACTAAGCAAGCAC) which is complementary to the 3'-noncoding region of the TSWV-BL N gene, and v (5'-TACAGTTCTAGAACCATGGATGATGCAAAGTCTGTGAGG) for the translatable of vi

(5'-AGATTCTCTAGACCATGGTGACTTGATGAGCAAAGTCTGTGAGGCTTGC)

for the untranslatablesecond half N gene fragment, the latter two oligomer primers are identical to the central region of the N gene. The oligomer primer lii contains a frameshift mutation immediately after the translation codon and several termination codons to block possible translation readthroughs while the oligomer primer vi contains several inframe termination codons immediately after the translation initiation codon.

The half gene fragments were purified on a 1.2% agarose gel as described above, and the gel-isolated gene fragments were digested with the restriction enzyme Ncol and directly cloned into Ncol -digested plant expression vector pBI525. The resluting plasmids were identified and designated as (1) pBI525-1N containing the first half translatable N gene, (2) pBI525-1N' containing the first half untranslatable N gene, (3) pBI525-1N- containing the first half translatable N gene in the antisense orientation, (4) pBI525-2N

- containing the second half translatable N gene, (5) pBI525-2n' containing the second half untranslatable N gene, and (6) pBI525-2N-containing the second half translatable N gene in the antisense ofientation. The expressin cassettes were then excised from plasmid pBI525 by digestion with *HindIII/EcoRI* and ligated as described above
- 35 into the plant transformation vector pBIN19 that had been cut with the

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same enzymes. The resulting vectors as well as plasmid pBIN19 were transferred to A. tumefaciens strain LBA4404, using the procedure described by Holsters supra. Leaf discs of N. benthamiana were inoculated with A. tumefaciens strain LBA4404 containing the various constructs. Transgenic plants were self-pollinated and seeds were self-pollinated and seeds were self-pollinated above.

Affalysis of transgenic plants by PCR and Northern hybridization PCR was performed on each R₀ transgenic line as described previously. The oligomer primers I to VI were used to determine the presence of the N coding sequence of TSWV-BL. The oligomer primer 35S-Promoter (see Example VIII) was combined with one of the above oligomer primers to confirm the orientation (relative to the CaMV 35S promoter) of the half gene sequences inserted into the plant genome. Northern analysis was conducted as described in Example VIII.

Lettuce isolate of TSWV (TSWV-BL) was used to challenge transgenic plants. Inoculation was done using test plants at the 3-4 leaf stage as described above. To avoid the possibility of escapes, control pants were used in each experiment and each inoculum extract was used to first inoculate the transgenic plants followed by control plants.

The various constructs used in this aspect of the present invention are illustrated in figure 8. Translatable and untranslatable half N gene fragemnts were synthesized by RT-PCR and then cloned directly into the plant expression vector pBI525. The oligomer primers ill and vi, used for generation of untranslatable half N gene fragments by RT-PCR, contains a mutation immediately after the translation initiation codon and the resulting reading frame contains several termination codons to block possible translation readthroughs. Thus, both first and second half untranslatable N gene fragments should be incapable of prodeing the truncated N protein fragments when introduced into plants. Both translatable and untranslatable half N gene fragements were then placed downstream of the CaMV 35S promoter of the vector pBI525 in the sense orientation or in the antisense orientation. The expressin of the half N coding sequences of TSWV-BL was thus controlled by a double CaMV 35S promoter fused to the 5'-

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untranslated leader sequence of alfalfa mosaic virus (ALMV) of the expression vector pBI525. Expression vectors that utilize the stacked double CaMV 35S promoter elements are known to yield higher levels of mRNA trnscription than similar vectors with a single 35S promoter element. Expression cassettes were transferred from the vector pBI525 to the pant transformation vector pBIN19. The resluting plasmids as well as the control plasmid pBIN19 were then transferred into A. tumefaciens strain LBA4404. Transgenic plants were obtained with nomenclature of the transgenic lines shown in figure 8.

All the kanamycin-resistant transgenic lines were confirmed by PCR to contain the proper N coding sequences in the expected orientations. Each transgenic R0 line which was grown for seeds was then assayed using Northern blot. Six out of six 1N, four out of six 1N', six out of six 1N-, six out of six 1N-, six out of six 2N, seven out of eight 2N', and six out of seven 2N- transgenic R0 lines were found to produce half N gene RNAs.

A set of transgenic R₀ plants was challenged with the homologous isolate TSWV-BL. Only asymptomatic plants were rated resistant while the plants showing any symptom (local lesions or systemic infections) were fated susceptible. All the inoculated R₀ control plants were susceptible to the virus; in contrast, two out of nine 1N', two out of six 1N-, four out of ten 2N', and one out of eight 2N-R₀ lines were found to be completely resistant to the virus infection. Although none of the 1N and 2N R₀ lines showed high levels of resistance, some of those plants displayed significant delays of symptom appearence.

Another set of transgenic Ro lines was brought to maturity for seed production. Seedlings were germinated on kanamycin-containing medium and inoculated with TSWV-BL. As shown in the following table, control seedlings and seedlings from some of the transgenic lines were susceptible to the isolate whereas seedlings from lines 1N-151, IN'-123, and 2N'-134 showed variojs levels of protection, ranging from delays in symptom expression to compete resistance.

	•	No. plant	s infected/No	. plants inoculated
	Ro line	6DPI	15DPI	30DPI
	Control	50/50		
	1N-149	17/17	_	
5	1N-151	2/20	13/20	17/20
	1N'-123	16/26	17/20	17/20
	1N'-124	20/20		
	· 1N'-126	19/19		
	1N=130	12/15	15/15	
10	1N~132	18/19	19/19	_
	2N-155	20/20		t v
	2N'-134	0/20	10/20	10/20
	2N'-135	19/19		
	2N ² -142	20/20		
15	2N143	20/20		
15	2N'*134 2N'*135 2N'-142	0/20 19/19 20/20	10/20	

In the above table, 30-fold diluted extracts of infected *N. benthamiana* were used to inoculate transgenic plants at the 3-4 leaf stage followed by control transgenic plants. DPI = days post inoculation.

In summary, this aspect of the present invention shows that transgenic plants expressing the first or the second half of either translatable or untranslatable N gene fragment are highly resistant to the homologous TSWV-BL isolate. This result demonstrates that a portion of the N gene is sufficient for resistance to the virus.

A listing of all nucleotide and amino acid sequences described in the foregoing description of the present invention is as follows:

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: Dennis Gonsalves and Sheng-Zhi Pang
 - (ii) TITLE OF INVENTION: Tomato Spotted Wilt Virus
- 3 0 (iii) NUMBER OF SEQUENCES: 30
 - (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH:

25 base pairs

(B) TYPE:

nucleic acid

3 5 (C) STRANDEDNESS: (D) TOPOLOGY:

single

(=) (=)

linear

(ii) MOLECULE TYPE:

DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

	ÁGCÁGGCÁÁÁ ÁCTOGCAGAA CITIGO 25	
	(2) INFORMATION FOR SEQ ID NO:2:	
	(i) SEQUENCE CHARACTERISTICS:	
ے	(A) LENGTH: 25 base pairs	
5	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(D) TOPOLOGY: tinëar (ii) MOLÉGULÉ TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
10	GCAAGITCIG CGAGITTIGC CIGCT 25	•
	(2) INFORMATION FOR SEQ ID NO:3:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 32 base pairs	
15	(B) TYPE: nucleic acid	
1 3	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLÉCULE TYPE: DNA	
	(xi) ŠEQUENCE DESCRIPTION: SEQ ID NO:3:	
$\hat{\mathcal{H}}_{i}$		
20	AGCTAÁCCÁT GGTTAAGCTC ACTAAGGAAA GC 32 (2) INFORMATION FOR SEQ ID NO:4:	
20	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 32 base pairs	
	(B) TYPE: nucleic acid	
2.5	(C) STRANDEDNESS: single	
25	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
	AGCATTCCAT GGITAACACA CIAAGCAAGC AC 32	
4.0	(2) INFORMATION FOR SEQ ID NO:5:	
30	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 2265 base pairs (B) TYPE: nucleic acid	
	(B) TYPE: nucleic acid (C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
35	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
	CAAGITGAAA GCAACAACAG AACIGIAAAT TCICITGCAG TGAAATCICT	50
	GCICATGICA GCAGAAAACA ACATCATGCC TAACTCTCAA GCTTCCACTG	100
	ATTCTCATTT CAAGCTGAGC CTCTGGCTAA GGGTTTTGAAG	150

	CAGGITICCA TICAGAAATT GITCAAGGIT GCAGGAGATG AAACAAACAA	200
•	AACATTTTAT TTATCTATTG CCTGCATTCC AAACCATAAC AGTGTTGAGA	250
	CĂĠĊĬĬĨĬĀĀĀ CĂĨĪTĀĊĨĠĨĬĪ ĂĨĬĬĬĠCĀĀGC ĀĬCĀGĊĪCCC ĀĀTĪCGCĀĀĀ	300
	TGCAAAGCTC CTTTTGAATT ATCAATGATG TTTTCTGATT TAAAGGAGCC	350
5	TTACAACATT GITCATGACC CITICATACCC CAAAGGATCG GITCCAATGC	400
	TCTGGGTCGA AACTCACACA TCTTTGCACA AGTTCTTTGC AACTAACTTG	450
	CAÁGÁAGATG TAÁTCATCTÁ CÁCTTTGÁAC AACCTTGÁGC TÁACTCCIGG	500
	AAAGTTAGAT TTAGGTGAAA GAACCITGAA TIACAGTGAA GATGCCTACA	550,
	AAAGCAAATÁ TTTCCTTTCA AAAACACTTG AATGTCTTCC ATCTAACACA	600
10	CAAÁCTÁTGT CTTACTTAGA CÁGCÁTCCAA ÁTCCCTTCAT GGAAGATAGA	650
	CITTOCCAGA GGAGAAATTA AAATTTCTCC ACAATCTATT TCAGITGCAA	700
	AATCTTTGTT AAAGCTTGAT TTAAGCCGGA TCAAAAAGAA AGAATCTAAG	750
	GITAAGGAAG CGIATGCTTC ÄGGATCAAAA TAATCITGCT TIGICCAGCT	800
	THTTCTAATT AIGHAIGH TATITICHT CHTACHAT AATTATTICH	850
15	CIGITITICIA TCICITICAÀ ÀTICCICCIG TCIAGIAGAA ACCATAAAAA	900
	CÁAÁÁÁTAA AAÁTGAÁÁAT ÁÁÁÁTTAAÁA TAAAATAAAA	1000
Ży.	ÀÀÀTÃÃÃAAC ÀÀCAÀAÁAT TÄÄAAAACGA AAAACCAAAA ÂGACCCGAAA	1050
	GGGACCAATT TGGCCAAATT TGGGTTTTGT TTTTGTTTTT TGTTTTTTGT	1100
	TTTTATTTT ATTTTATTT TATTTTATTTTTTTTTTT	1150
20	ATTTATTIA TTITTIGITT TOGITGITTT TGITATTIA TIATTIATTA	1200
	ÀGCĂCĂACAC ACĂGAAÁGCA ÁACTITAATT AAACACACTT ÁTITAAAATT	1250
	TAÁCÁCACTA AGCAAGCÁCÁ ÁGCAATAAAG ATAAAGAAAG CITTATATAT	1300
	THATAGCTT TTTTATAATT TAACTTACAG CIGCTTTCAA GCAAGTTCTG	1350
	CGAGTITIGC CIGCITTITA ACCCCGAACA TITCATAGAA CITGITAAGA	1400
25	GITTCACIGI AATGITCCAT AGCAACACIC CCITTAGCAT TAGGATTGCT	1450
	GCAGCTAAGT ATAGCAGCAT ACTCTTTCCC CTTCTTCACC TGATCTTCAT	1500
	TCÁTTICAAA TGCTTTGCTT TTCAGCACAG TGCAAACTTT TCCTAAGGCT	1550
	TOCTTGGIGT CATACTTCTT TGGGTCGATC CCGAGGTCCT TGTATTTTGC	1600
	ATCCTGATAT ATAGCCAAGA CAACACTGAT CATCTCAAAG CTATCAACTG	1650
30	AAGCÁÁTAAG AGGTAAGCTA CCTCCCAGCA TTATGGCAAG TCTCACAGAC	1700
	TTTGCÁTCAT CGAGAGGTÁA TCCATAGGCT TGAATCAAAG GÁTGGGAÁGC	1750
	AATCTTAGAT TIGATAGIAT TGAGATTCIC AGAATICCCA GITTCITCAA	1800
	CAAGCCIGAC CCIGATCAAG CIATCAAGCC TICTGAAGGI CATGICAGIG	1850
	CCICCAÀICC IGICIGAAGI TITICITIAIG GIAATITIAC CAAAAGIAAA	1900
35	ATCGCTTIGC TTAATAACCT TCATTATGCT CTGACGATIC TTTAGGAATG	1950

	TCAGACATGA AATAACGCIC ATCITCITGA TCIGGICGAT GITTTCCAGA	2000
	CĂĂĂĂAGICT TGĂĂĞTTGĂĂ TGCTACCĂGĂ TICTGATCIT CCTCAAACTC	2050
	AAGGICITIG CCITIGIGICA ACAAAGCAAC AATGCTTTCC TTAGTGAGCT	2100
	TĂĂČĆĪTĀGĀ CATGĀĪGĀĪĆ GTĀAĀĀGITG TTATĀGČĪTT GĀCCGTATGT	2150
5	AÁCTCAÁGGT GOGAÁÁGTGC ÁÁCTCTGTÁT COCGCÁGTCG TTTCTTAGGT	2200
	TCTTAATGTG ATGATTTGTA AGACTGAGTG TTAACGTATG AACACAAAAT	2250
	TGACACGAÍTT GCTCÍ 2265	
	(2) INFORMATION FOR SEQ ID NO:6:	,
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1709 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	·
	(II) MOLECULE TYPE: DNA	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
	AAATTCTCTT GCAGTGÁÁÁT CTCTGCTCAT GTTAGCAGAA AACAACATCA	50
	TGCCTAACTC TCAAGCTTTT GTCAAAGCTT CTACTGATTC TAATTTCAAG	100
	CTGAGCCTCT GGCTAAGGGT TCCAAAGGTT TTGAAGCAGA TTTCCATTCA	150
	GAAATTGTTC AAGGITGCAG GAGATGAAAC AAATAAAACA TTTTATTTAT	200
20	CTATTGCCTG CATTCCAAAC CATAACAGTG TTGAGACAGC TTTAAACATT	250
	ACTIGITATITI GCAAGCATCA GCTCCCAATT CGTAAATGTA AAACTCCTTT	300
	TGÁÁTTATCA ÁTGATGITTT CIGATTIAAA GGAGCCTTAC AACATTATTC	350
	ATGATCCTTC ATATCCCCAA AGGATTGTTC ATGCTCTGCT TGAAACTCAC	400
	ACATCITTIG CACAAGITCT TIGCAACAAC TIGCAAGAAG ATGIGATCAT	450
25	CTÁCÁCCTTG ĂACÁACCATG ÁGCTAACTCC TGGAAAGTTA GATTTAGGTG	500
	AÁÁŤAACTÍT GAÁTÍTÁCÁÁT GÁAGACGCCT ACAAAAGGAA ATATTTCCTT	550
	TCAAAAACAC TTGAATGTCT TCCATCIAAC ATACAAACTA TGTCTTATTT	600
	AGÁCÁGCATC CAAÁTOCCTT OCTOGAAGAT AGACTITICCC AGOGGAGAAA	650
	TTAAAATTIC TCCACAATCT ATTTCAGITG CAAAATCTTT GITAAATCTT	700
30	GATTTAAGCG GGATTAAAAA GAAAGAATCT AAGATTAAGG AAGCATATGC	750
	TICÁGGÁTCÁ ÁAATGATCIT GCIGIGICCA GCITTITCIA ATTATGITAT	800
	GITTATTTIC TETCTTACT TATAATTATT TETCTGTTIG TCATTICTTT	850
	CAAATTCCTC CTGTCTAGTA GAAACCATAA AAACAAAAAT AAAAATAAAA	900
	ΤΆΑΑΑΤΚΑΑΑ ΑΤΑΑΑΑΤΑΑΑ ΑΤΑΑΑΑΤΑΑΑΑ ΤΟΑΑΑΤΑΑΑΑ ΘΟΑΑΟΑΑΑΑΑ	950
3 5	AATTAAAAAA CAAAAAAACCA AAAAAGATCC CGAAAGGACA ATTITIGGCCA	1000
	AATTIGGGT TIGHTTIGH TITHGHTT TITGHTTIT GHTHTATTT	1050

	THATHTHAT TITHATHTIT ANTHATTIT ATTHATGIT THIGHTGIT 11	L00
	TTGTTÄTTTT GTTATTTATT ÅÅGCACAACA CACAGAAAGCA AACTTTAÁT 1	L50
	TAAACACACT TATTTAAAAT TTAACACACT AAGCAAGCACA AACAATAAA 1	200
	GÁTÁAÁGÁAA GCTTTÁTÁTA TTTATÁGGCT TTTTTATÁAT TÍTÁACTTÁCA 1	250
5	CCTCCTTTTA ACCAACITICT CICAGITTTG CCTGTTTTTT AACCCCAAAC 1:	300
	ATTICATAGA ACTIGITAAG GETTICACIG TAATGITICA TAGCAATACT 1:	350
	TOCTTTÁGCA TTAGGÁTTGC TGGÁGCTAAG TATAGCÁGCÁ TÁCTCTTTCC 1	400
	CCTTCTTCAC CTCATCTTCA TTCATCTCAA ATGCTTTTCT TTTCAGCACA 1	450
	GÜGCĂĂĂCIT TÜCCTAAGGC TÍCCCTGGIG TCATACTICT TTGGGICGAT 1	500
10	COCGAGATCC TIGIATTITIG CATOCIGATA TATAGCCAAG ACAACACIGA 1	550
	TCATCTCAAA GCTATCAACT GAAGCAATAA GAGGTAAGCT ACCTCCCAGC 1	600
	ÁTTÁTGGCÁA GCCTCÁCÁGÁ CTTTGCÁTCA TCAAGÁGGIA ATCCATAGGC 1	650
	TIGÁATCAAA GGGIGGGÁAG CÁATCTIAGA TITGATAGIA TIGAGATICT 1	700
	CAGAATTCC 1709	
15	(2) INFORMATION FOR SEQ ID NO:7:	
	(I) SEQUÊNCE CHARACTERISTICS:	
$P_{\mathcal{F}}$	(A) LENGTH: 260 amino acids (B) TYPE: amino acid	
	(C) STRANDEDNESS: single	
20	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: peptide	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
	Gln Val Glu Ser Åsn Asn Arg Thr Val Asn Ser Leu Ala Val 5 10	Lys 15
25	Ser Leu Leu Met Ser Ala Glu Asn Asn Ile Met Pro Asn Ser	
	20 25	30
	Ala Ser Thr Asp Sér His Phe Lys Leu Ser Leu Trp Leu Arg 35 40	45
	Pro Lys Val Leu Lys Gln Val Ser Ile Gln Lys Leu Phe Lys	
30	50 55	60
	Ala Gly Asp Glu Thr Asn Lys Thr Phe Tyr Leu Ser Ile Ala 65 70	75
	Ile Pro Asn His Asn Ser Val Glu Thr Ala Leu Asn Ile Thr	
äE	80 85	90
35	Ile Cys Lys His Gln Leu Pro Ile Arg Lys Cys Lys Ala Pro 95 100	105
	Glu Leu Ser Met Met Phe Ser Asp Leu Lys Glu Pro Tyr Asn	
	110 115	120
40	Val His Asp Pro Ser Tyr Pro Lys Gly Ser Val Pro Met Leu 125 130	_
. 0	120 100	135

,	Leu Glu	Thr	His	Thr 140	Ser	Leu	His	Lys	Phe 145	Phe	Ala	Thr	Asn	Leu 150
	Gln Glu	Ásp	Val		11e	Tyr	Thr	Leu		Asn	Leu	Glu	Leu	
5	Pro Gly	Lys	Leu		Leu	Gly	Glu	Arg		Leu	Ásn '	Tyr	Ser	
	Asp Ala	Tyr	Lys		Asp	Tyr	Phe	Leu		Lys	Thr	Leu	Glu	
10	Leu Pro	Ser	Asn	Thr 200	Gl'n	Thr	Met	Ser	Tyr 205	Léu	Asp	Ser	Ile	Gln 210
	Ile Pro	Ser	Trp	Lys 215	Ile	Asp	Phe	Ala	Arg 220		Glu	Ile	Lys	Ile' 225
	Ser Pro			230					235					240
15	Leu Ser	Gly	Ile	Lys 245	Lys	Lys	Glu	Ser	Lys 250		Lys	Glu	Ala	Tyr 255
	Ala Ser	Gly	Ser	Lys 260										
	(2) INFO	RMAT	ΓΙΟN	FOR	SEQ	ID N	:8:C							
20	(i)	SEC		CEC			RIST	ICS:						
			` '	ENG					e pa	irs				
			` '	YPE:		NES			acid ingle	ı				
			` '				o. Iine		ყ.					
25	(i	i) MO	` '	JLE T			DN	4						
	()	d) SE	QUE	NCE	DESC	RIPT	TON:	SEQ	ID N	O:8:				
	TTAACAC	CACT	AAG	ZAAGO	AC A	VÁACA	ATAP	A G	SAAT!	GAAZ	GC1	'I'I'A	ATA	50
	TTTATA	GCT	TTT	LTATZ	L TA	TAAC	TTAC	A GO	CIGCI	1117	A AGO	[AAG]	TCT	100
	GIGAGI.	ITIG	CCI	311 ¹ 11	ÀT <i>i</i>	AACCC	CAA	AC AT	rtrcz	ATTAG?	A ACI	rigr	CAAG	150
30	GGITIC	ACIG	TAAT	IGII	CA!	L'AGC2	ATA/	CT TO	CIT	L'AGC	A TIZ	AGGAT	rigc	200
	TGGAGC	TAAG	TAT	AGCA(CÀ!	IACI	CTTC	cc a	CIIC	TCA(CIY	AIC.	ITCA	250
	TICATT	ICAA	ATG	CITT.	ICI :	rric	AGCA(CA G	IGCAZ	AACT.	r TI	CIA	AGGC	300
	TTCCCT	GGIG	TCA'	IACI'	ICI '	IIGG	FICE/	AT C	CCGA	CATC	CTI	GLAT.	ITTG	350
	CATCCI	GATÁ	TAT	AGCC2	AAG I	ACAA	CACIV	GA TY	CATC	ICAA	A GC	IATC	AACT	400
3 5	GAAGCA	ATTA	GÀG	GTAA	GCT I	ACCIV	CCCA	GC A	TATI	GCA)	A GC	CICA	CAGA	
	CITIGO													
	CAATCT	TAGA	ΪΠ	GATA	GIÀ '	TIGA	GATT	CT C	AGAA'	TTCC	C AG	TTTO	CTCA	~ 5 50
	ACAAGO	CTGA	ccc	TGĀT	CAÁ	GCTA'	ICAA	GC C	TICI	GAAG	G TC	AIGI	CAGI	600
	GGCICC													
4.0	רביידועע	יעודווי	C TTT	a á má	X CYC 1	ותריעונוו	ידו לינדר	~~ m	עניאוד)	ייווערייי	T (M)	תיא ה	~ n m	700

GICAGACATG AAATAATGCT CATCITTTIG ATCIGGICAA GGITTICCAG	750
ACÁAÁÁAGIC TTGÁÁGTTGÁ ÁTGCIÁCCAG ÁTTCTGÁTCT TOCTCAAÁCT	800
CĂĂĞĞİCTIĞ GCCÜĞÜĞÜC AACAAAGCAA CAATGCTÜÜC CÜTAGIGAGC	850
TTÄÄÕČAT 858	
(2) INFORMATION FOR SEQ ID NO:9:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 2028 base pairs	
(B) TYPE: nucleic acid (C) STRANDEDNESS: single	,
(D) TOPOLOGY: linear	·
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
AAATTICTCTT GCAGTGAAAT CTCTGCTCAT GTTAGCAGAA AACAACATCA	50
TGCCTÂACTC TCAAGCTTTT GICAAAGCTT CTACTGATTC TAATTICAAG	100
CTGAĞCCICT GGCIÁAGGĞI TCCAAAGGIT TIGAAGCAGA TITCCATICA	150
GAÂÁÍÍÍGTÍC AAGGTÍGCÁG GAGÁTGAÁAC AAATAAAACA TTTTÁTTTAT	200
CTÁTTGCCTG CÁTTCCÁÁÁC CÁTÁACAGTG TTGAGACAGC TTTÁAACÁTT	250
ACIGITATTT GCAAGCATCA GCTCCCAATT CGTAAATGTA AAACTCCTTT	300
TGAĂȚIATCĂ ATGATGITIT CIGATITAAA GGAGCCTIAC AACATTATIC	350
ATGATCCITC ATATCCCCAA AGGATIGITC ATGCTCTGCT TGAAACTCAC	400
ÁCĂÍCHTTG CÁCAÁGITCT TIGCAACAAC TIGCAAGAAG ATGIGATCAT	450
CIÁCACCITG AACAACCATG AGCTAACICC TGGAAAGITA GATTIAGGIG	500
AAATAACITT GAATTACAAT GAAGACGCCT ACAAAAGGAA ATATTICCIT	550
TCAAAAACAC TTGAATGICT TCCATCIAAC ATACAAACIA TGICITATIT	600
AGACÁGCAÍC CAAAICCCÍÍ CCIGGAAGAT AGACITIGCC AGGGGAGAAA	650
TTÄÄÄÄTTIC TCCACAÄTCT ATTICAGITG CAAAATCITT GITAAATCIT	700
GATTTAAGCG GGATTAAAAA GAAAGAATCT AAGATTAAGG AAGCATATGC	750
TICAGGATCA AAATGATCTT GCTGTGTCCA GCTTTTTCTA ATTATGTTAT	
GITTATTIC TITCITTACT TATAATTATT TITCIGITIG TCATTICITT	
CAÀÁTTCCTC CTGTCTÁGTÁ GAAACCATAA AAACAAAAAT AAAAATAAAA	900
TAÁAÁÍCAAA ÁTAAÁATAAA ÁATCAAAAAA TGAAATAAAA GCAACAAAAA	
AÁÍTTÁÁAAÁÁ CAAAÁAACCA AAÁAAGATCC CGAAAGGACA ÁTTTTGGCCA	1000
AATTIGGGT TIGITTTIGT TTTTTTTTTTTTTTTTTTT	`
TTÄTTTTAT TTTTATTTT ATTTATTTT ATTTTATGIT TTTGTTGTT	1100
TIGITATITI GITATITATI AAGCACAACA CACAGAAAGC AAACTITAAT	
TAAACACACT TATTTAAAAT TTAACACACT AAGCAAGCAC AAACAATAAA	1200

	GAŤAÂAGAAA GCTTTATATA TTTATAGCCT TTTTTATAAT TTAACITACA	1250
	GCTGCTTTTA ÀGCAAGTTCT GTGAGTTTTG CCTGTTTTTT ĂĂCCCCAAAC	1300
	ATTICATAGA ACTIGITAAG GGITTCACIG TAATGITOCA TAGCAATACT	1350
	TOTTITAGCÁ TIAGGÁTIGO TIGGÁGOTAÁG TATÁGCAGCÁ TÁCTOTTICO	1400
5	CCTTCTTCAC CTGATCTTCA TTCATTTCAA ATGCTTTTCT TTTCAGCACA	1450
	GIĞÜAAACIT TÜCCIAAGGC TÜCCCİGGIG TCATACTÜCÜ TÜĞGĞİCGAT	1500
	CCCGAGATCC TTGIATTTTG CATCCTGATA TATACCCAAG ACAACACTGA	1550
	TCÁTCTCAAA GCTATCAACT GAAGCAATAA GAGGTAAGCT ACCTCCCAGC	1600,
	ÁTTATGGCÁÁ GÖCTCÁCÁGÁ CTTTGCÁTCÁ TCÁAGÁGGTA ÁTCCÁTAGGC	1650
10	TTGÁCTCÁAÁ GGGTGGGÁÁG CAÁTCTTÁGA TTTGATÁGTÁ TTGÁGATTCT	1700
	CAGAATICCC AGITICCICA ACAAGCCIGA CCCTGATCAA GCTATCAAGC	1750
	CITCIGAAGG TCATGTCAGT GOCTOCAATC CIGICIGAAG TTTICITTAT	1800
	GGTÄÄTTITA CCAAAAGTÄA ÄATCGCTITG CTTAATAACC TTCATTATGC	1850
	TCTGACGATT CTTCAGGAAT GTCAGACATG AAATAATGCT CATCTTTTTG	1900
15	ATCTGCTCAA GGTTTTCCAG ÁCAAAAAGIC TIGAAGITGA ÁTGCTACCAG	1950
	ATTOTOATOT TOCTOANACT CAAGGICTTT GCCTTGTGTC AACAAAGCAA	2000
1	CAATGCTTTC CTTAGTGAGC TTAACCAT 2028	
	(2) INFORMATION FOR SEQ ID NO:10:	
4.0	(i) SÉQUENCE CHARACTERISTICS:	
20	(Á) LENGTH: 22 base pairs (B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(II) MOLECULE TYPE: DNA	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
	TICIGGICIT CITCAAACT CA 22	
	(2) INFORMATION FOR SEQ ID NO:11:	
	(i) SEQUÊNCE CHARACTERISTICS:	
	(A) LENGTH: 18 base pairs	
.30	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(II) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	` `
35	(XI) SEQUENCE DESCRIPTION. SEQ 10 NO. 11: CIGIÁGOCAT GÁGCAAAG 18	•
55	(2) INFÓRMATION FOR SEQ ID NO:12:	
	(i) SEQUENCE CHARACTERISTICS:	
	WOLKOLIYOL OF MINOTENIO 1100.	

			((B) Th	HAN	DEDI		ämir :	no ad sil	no a cid nglë	cids				
5		2111				OGY:		line							
Э		• •			LETY		بند عرب	pepi		· 6 4	. 1 =				
e.	4.4-1					EŠČI									
	Met	ser	ser	GTĀ	Val 5	Tyr	Glu	Ser	Ile	Ile 10	Gln	Thr	Lys	Ala	Ser 15
10	Val	Trp	Gly	Ser	Thr 20	Alä	Ser	Gly	Lys	Ser 25	Ile	Val	Asp	Ser	Tyr 30
	Trp	Ile	Tyr	Glu	Phe 35	Pro	Thr	Gly	Ser	Pro 40	Leu	,Val	Gln	Thr	Gln 45
	Leu	Tyr	Ser	Asp	Ser 50	Arg	Ser	Lys	Ser		Phe	Gly	Tyr	Thr	
15	Lys	Ile	Gly	Åsp	Ile 65	Pro	Ala	Val	Glu		Glu	Ile	Leu	Ser	
	Asn	Val	His	Ile	Pro 80	Val	Phe	Asp	Asp	Ile 85	Asp	Phe	Ser	Ile	Asn 90
,20	Ile	Ash	Asp	Ser	Phe 95	Leu	Ala	Ile	Ser	Val 100	Cys	Ser	Asn	Thr	
,	Asn	Thr	Asn	Gly	Val 110	Lys	His	Gln	Gly	His 115	Leu	Lys	Val	Leu	Ser 120
	Leu	Alá	Gln	Ĺeu	His 125	Pro	Phe	Glu	Pro		Met	Ser	Arg	Ser	
25	Ile	Åla	Ser	Arg	Phe 140	Arg	Leu	Gln	Glu		Asp	Ile	Ile	Pro	Asp 150
	Asp	Lys	Tyr	Ile	Ser 155	Ala	Ala	Asn	Lys		Ser	Leu	Ser	Cys	Val 165
30	Lys	Glu	His	Thr	Tyr 170	Lys	Val	Glu	Met		His	Asn	Gln	Ala	Leu 180
	Gly	Lys	Val	Àsn	Val 185	Leu	Ser	Pro	Asn	-	Asn	Val	His	Glu	
	Leu	Tyr	Ser	Phe		Pro	Asn	Glu	Asn		Ile	Glu	Ser	Asn	
35	Arg	Thr	Val	Asn	Ser 215	Leu	Ala	Val	Lys		Leu	Leu	Met	Ala	
	Glu	Asn	Asn	Ile		Pro	Asn	Ser	Gln		Phe	Val	Lys	Ala	Ser 240
40	Thr	Ásp	Ser	His		Lys	Leu	Ser	Leu		Leu	Arg	Ile	Pro	Lys 255
	Val	Leu	Lys	Gln		Ala	Ile	Gln	Lys	Leu 265	Phe	Lys	Phe	Ala	Gly 270
	Asp	Glu	Thr	Gly		Ser	Phe	Tyr	Leu		Ile	Àla	Cys	Ile	Pro 285

	Asn	His	Asn	Ser	Val 290	Glu	Thr	Ala	Leu	Asn 295	Val	Thr	Val	Ile	Cys 300
	Arg	His	Gln	Leu		Ilė	Pro	Lys	Ser	-	Ala	Pro	Phe	Glu	
5	Ser	Met	Ilė	Phe		Ásp	Leu	Lys	Glu		Tyr	Ásn	Thr	Val	
	Asp	Pro	Ser	Tyr	Pro 335	Gĺń	Arg	Ile	Val		Ala	Leu	Leu	Glu	
10	His	Thr	Ser	Phe	Å 1á 350	Gln	Val	Leu	Cys	Asn 355	Lys	Leu	Gln	Glu	Asp 360
	Val	Ilé	Ile	Tyr	Thr 365	Ile	Asn	Ser	Pro	Glu 370	Leu	Thr	Pro	Ala	
	Leu	Àśp	Leu	Gly	Glu 380	Arg	Thr	Leu	Asn	Tyr 385	Ser	Glu	Asp	Ala	
15	Lys	Lys	Lys	Tyr	Phe 395	Leu	Ser	Lys	Thr	Leu 400	Glu	Cys	Leu	Pro	Val 405
					410					415			Ile		420
20					425					430			Ser		435
7					440					445			Leu		450
0 F				Lys	Lys 45 5	Ser	Leu	Thr	Trp	Glu 460		Ser	Ser	Tyr	Asp 465
25	Leu	Glu	;												
	(2) !	NFOI	TAMF	TION	FOR	SEQ	ID N	O:13:							
							ACTE								
		(7					.0.2		ami	no s	cids				
					YPE:				no a						
30				(c) s	MART	NDED	NES:			ingle					
				(D) T	opol	OGY	:	line	ar						
		(ii) MOI	LECL	ILET	YPE:		per	tide						
		(x	i) SE	QUEN	ICE t	DESC	RIPT	ION:	SEQ	ID N	O:13:				
35	Met	Ser	Lys	Val	Lys 5		Thr	Lys	Glu	Asn 10		Val	. Ser	Leu	Leu 15
	Thr	Gln	Ser	Ala	Asp 20		Glu	Phe	Glu		Asp	Glr	Asn	Gln	
	Ala	Phe	Asn	Phe	Lys 35		Phe	Cys	Glr		Asn	Let	Asp	Leu	
40	Lys	Lys	Met	. Ser	: Ile 50		Ser	Cys	Leu		Phe	Leu	ı Lys	Asn	
	Gln	Gly	' Ile	Met	Lys 65		. Val	. Asn	Glr		Asp	Phe	Thr	Phe	

	Lyš V	al	Thr	İle	Lys 80	Ĺýŝ	Asn	Ser	Glu	Arg 85	Val	Gly	Ala	Lys	Asp 90
	Met T	hr	Phe	Àrg	Arg 95	Leu	Åsp	Ser	Met	Ile 100	Arg	Val	Lys	Leu	
5	Glu Ġ	lu	Thr	Åla	Ash 110	Āšn	Glu	Àsn	Leu	Ala 115	Ile	Ile	Lys	Ala	Lys 120
	Ílë Å	la	Sér	His	Pro 125	Léu	Val	Gln	Ala	Tyr 130	Gly	Leu	Pro	Leu	
10	Åsp Å	ĺa	Lys	Ser	Val 140	Arg	Leu	Ala	Ile	Met 145	Leu	Gly	Gly	Ser	
•	Pro I	Ĕu	Ile	Àla	Ser 155	Val	Asp	Ser	Phe	Glu 160	Met	Ile	Ser	Val	
	Leu A	la	Ile	Tyr	Gln 170	Asp	Ala	Lys	Tyr	Lys 175	Glu	Lėu	Gly	Ile	
15	Pro 1	hr	Lys	Tyr	Asn 185		Lys	Glu	Ala	Leu 190	Gly	Lys	Val	Cys	Thr 195
	Val I	eu	Lys	Ser	Lys 200		Phe	Thr	Met	Asp 205		Ala	Gln	Ile	Asn 210
20	Lys (Зlý	Lys	Glu	Tyr 215		Lys	Ile	Leu	Ser 220	Ser	Cys	Asn	Pro	Asn 225
$\hat{\mathcal{H}}$	Åla I	<u>i</u> ys	Gly	Ser	Ile 230		. Met	Asp	Tyr	Tyr 235		Asp) Asn	Leu	Asp 240
	Lys I	Phe	Tyr	Glu	Met 245		Gly	Val	Lys	Lys 250		Ala	Lys	Ile	Ala 255
25	Glý (/al	Ala												
	(2) IN	FÖI	AMF	TION	FOR	SEC	ID N	O:14	:						
		(i)	SEG	UEN	CEC	HAR	ÅCTE	RIST	TCS:						
				(Å) L	ENG	TH:			9 ba	ase p	airs				
4.0				` '	YPE:		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \		eleic						
30				` '		LOGY	NES ,.	S: line		ingle	!				
		/::	MA												
		•	•				CRIPT		-	in ki	044				
	AGAG	•	•										\ ~~ ~1\ 7	70 m/C	5 0
35	TCÁG														50 100
55	TTÌÀ														
	TCİĞ									^					200
	TICH													_	
							ATTG								300
40	ATTT														350
							ÍCTI								400

	TTÁÁCÁCCAA TOGÁG	TGÁÁG CÁTCÁGGGIC	ATCTTAAAGT	TCTTTCTCTT	450
		THICA ACCIGIGATE			500
	_	ŘĠŘŘG ŘŘGÁTÁŤÁÁŤ			550
	CIGCIGCIAA CAAGG	GATCT CICICCIGIO	TCÁÁÁGÁÁCÁ	TACTTACAAA	600
5	GICGAAATGA GCCAC	AATCA GGCTTTAGGC	AAAGÏGĀĀĪ G	TICITICICC	650
	TAACAGAAAT GITCA	iigagii gciigiatac	TTTCAÂACCA	ÂĂTTICAACC	700
•	ĂĠĂĨŎĠĂĂĀG ŤÁĂŤĀ	ācāgā ācigiaaāti	CICTIGCAGT	CAAATCTTTG	750
		AACAA CATTATGCCI			800
•	AGCITCIACT GATTO	TCATT TTAAGITGAG	CCITIGGCIG	ÁGAÁTTCCAA	850 [°]
10	AAĞITTIĞAA GCAAA	TAGCC ATACAGAAGC	TCTTCAAGIT	TÇCAGGAGAC	900
		TICIA TIIGICIATI			950
		TITAA ATGICACIGI			1000
٠	CAATOCCIAA GICCA	AAGCT CCTTTTGAAT	TATCAATGAT	TITCICCGAT	1050
	CIGAAAGAGC CITAC	AACAC TGTGCATGAT	CCTTCATATC	CTCAAAGGAT	1100
15	TGITCATGCT TIGCT	TGÅGA CTCACACTTC	CITIGCACAA	GITCICIGCA	1150
	ACAAGCTGCA AGAAG	ATGIG ATCATATATA	CTATAAACAG	CCCTGAACTA	1200
<i>?</i>	ACCCCÁGCTA AGCTO	GATCT ÁGGTGAAÁGA	ACCITGAACT	ÀCAGTGAAGA	1250
•	TGCTTCGAAG AAGAA	GIÁTT TICTTICAA	AACACICGAA	TGCTTGCCAG	1300
		TGICT TATTIGGATA			1350
20	AAGÁTAGACT TTGCC	AGAGG AGAGAICAG	ATCTCCCCTC	ÁATCTACTCC	1400
	TATTICCAAGA TCTTT	GCICÁ ÀGCIGGATIT	GAGCAAGATC	AAGGAAAAGA	1450
	AGICCITGAC TIGGO	AAACA TOCAGOTATO	ÀTCIAGAATA	ÁÁAGIGGCIC	1500
	ATACIACICI AAGTA	GIÁTT TGÍCAACTT	CTTATCCTTT	ATGITGITTA	1550
	TTTCTTTTAA ATCTA	AAGIA AGITAGATIY	AAGIAGIITA	GIATGCIATA	1600
25	GCATTATTAC AAAAA	ATÁCA ÁAAAAATAC	AAAAAATACA	AATATAAÁA .	1650
	AAAACCCAAA AAGAT	CCCAA AAGGGACGA	TIGGIIGATT	TACTCIGITT	1700
	TAGGCTTATC TAAGC	CIGCIT TIGITIGAG	AAAATAACAT	TGTAACATGC	1750
	AATAACTGGA ATTTA	VAAGIC CIAAAAGAA	G TTTCAAAGGA	CAGCTTAGCC	1800
	AAAATIGGIT TIIGI	TITIE TITITE	TITIGITIT	TIGITTTÄTT	1850
30	TITATTITTA GITT	ATTIT TGFFFFFGF	TTTATTTATTT	TTATTTTATT	1900
	TICITITATI TIATI	TATAT ATATATCAA	A CACAATCCAC	ÁCAAATAATT	1950
	TTAATTICAA ACAT	ICÍACT GATTIAACA	C ACITAGCCIG	ACITIATCAC	2000
	ACTTAACACG CTTAC	STEAGG CTTTAACAC	A CIGAACIGAA	TTAAAACÁCA	2050
	CITAGÜATTA TOCAT	CICIT AATTAACAC	A CTTTAATAAT	ATGCATCTCT	2100
3 5	GAATCAGCCT TAAAC	AAGCT TTTATGCAA	CACCAGCAATC	TIGGCCTCTT	2150

	TCTTAACTCC AAACATTTCA TAGAATTTGT CAAGATTATC ACTGTAATAG	2200
	TOCATAGCAA TECTICOCTI AGCATTEGGA TIGCAAGAAC TAAGIATCIT	2250
	GGCATATICT TICCCTITIGT TIATCTGTGC ATCATCCATT GTAAATCCTT	2300
	TOCTITIAAG CACTGIGCAA ACCITICCCA GAGCITCCTT AGTGTTGTAC	2350
5	THAGITIGGIT CAATCCCHAA CICCTIGHAC THIGCATCH GATATATGGC	2400
	AAGAACA CIGATCATCT CGAAGCTGTC AACAGAAGCA ATGAGAGGGA	2450
	TACTACTIC AAGCATTATA GCAAGICTCA CAGATTITIGC ATCTGCCAGA	2500
	GGCÁGCCCT AÁGCTTGGÁC CAÁÁGGCTGG GAGGCÁATTT TIGCTTTGAT	2550
	AATAGEAAGA TICTCATIGT TIGCAGTCIC TICTATGAGC TICACICITA	2600
10	TCATGCTATC AAGCCTCCTG AAAGTCATAT CCTTAGCTCC AACTCTTTCA	2650
	GAATTTTCT TTATOGIGAC CTTACCAAAA GTAAAATCAC TTTGGTTCAC	2700
	AACTITICATA ATGCCTTIGGC GATTCTTCAA GAAAGTCAAA CATGAAGTGA	2750
	TACTCATTTT CTTAATCAGG TCAAGATTTT CCTGACAGAA AGTCTTAAAG	2800
	TTGĂĂTGCGA CCTGGTTCTG GICTTCTTCA AACTCAACAT CTGCAGATTG	2850
15	AGTTAAAAGA GAGACAATGT TTTCTTTTGT GAGCTTGACC TTAGACATGG	2900
	TGGCAGTTTA GATCIAGACC TTTCTCGAGA GATAAGATTC AAGGTGAGAA	2950
	AGIGCAACAC TGIAGACCGC GGICGITACT TATCCIGITA ATGTGATGAT	3000
	TIGIATIGCT GAGIATTAGG TITTIGAATA AAATIGACAC AATIGCTCT	3049
	(2) ÎNFÓRMÁTION FOR SEQ ID NO:15:	
20	(I) SEOUENCE CHÁDACTEDISTICS:	

(A) LENGTH:

778 base pairs

(B) TYPE:

Nucleic acid

(C) STRANDEDNESS:

single

(D) TOPOLOGY:

linear

25 (ii) MOLECULE TYPE:

DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

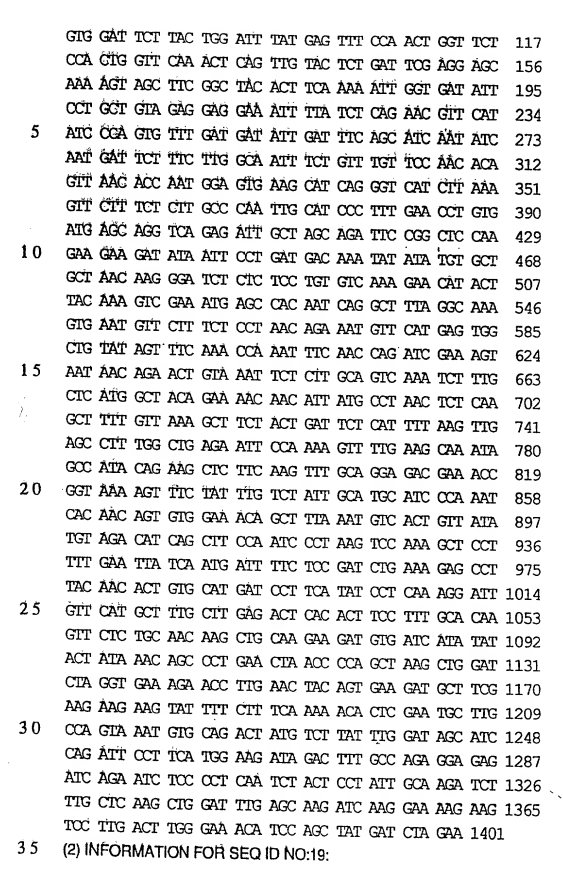
ATG CAA CAC CÂG CÂĂ TCT TGG CCT CTT TCT TAA CTC CAA 39 ACA TIT CAT AGA ATT TGT CAA GAT TAT CAC TGT AAT AGT 78 CCA TAG CAA TGC TTC CCT TAG CAT TGG GAT TGC AAG AAC 117 30 TAA GIA TCT TGG CAT ATT CIT TCC CIT TGT TTA TCT GIG 156 CAT CAT CCA TIG TAA AIC CIT TGC TTT TAA GCA CIG TGC 195 AAA CCT TCC CCA GAG CIT CCT TAG TGT TGT ACT TAG TTG 234 GIT CAA TCC CIA ACT CCT TGT ACT TTG CAT CIT GAT ATA 273 \ TGG CÁA GAÁ CAA CÁC TGÁ TCA TCT CGA AGC TGT CAÁ CÁG 312 35 AAG CAA TGA GAG GGA TAC TAC CTC CAA GCA TTA TAG CAA 351

GIC TCA CAG ATT TIG CAT CIG CCA GAG GCA GCC CGT AAG

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, CÎTÎ GGÂ CCA AAG GGT GGG AGG CAA TIT TIG CIT TGA TAA 429 TAG CAA GAT TĈI CAT TGI TIG CAG TCT CIT CIA TGA GCT 468 TCA CTC TIÁ TCÁ TGC TÁT CAA GOC TCC TGA ÁAG TCA TAT 507 CCT TAG CIC CAA CTC TTT CAG AAT TIT TCT TTA TCG TGA 546 CCT TAG CAA AAG TAA AAT CAC TIT GGT TCA CAA CIT TCA 585 TÀA TGC CIT GGC GÁT TCT TCA ÁGÁ ÁAG TCÁ AAC ÁTG AAG 624 TIGA TAC TOA TIT TOT TAA TOA GGT CAA GAT TIT COT GAC 663 AGA ÂAG TCT TAA ÂGT TGA ÂTG CGA CCT GGT TCT GGT CTT 702 CIT CĂĂ ACT CĂA CĂT CTG CAG ĂTT GAG TIA ÁAA GAG AGA 741 10 CAÁ TGT TIT CIT TTG TGA GCT TGA CCT TAG ACA TGG (2) INFORMATION FOR SEQ ID NO:16: (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: Nucleic acid 15 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA 17 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16: GITCIGAGAT TIGCTAGT 18 (2) INFORMATION FOR SEQ ID NO:17: 20 (I) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: Nucleic acid (C) STRANDEDNESS: single 25 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17: TIATATCITC TICITGGA 18 (2) INFORMATION FOR SEQ ID NO:18: 30 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1401 base pairs (B) TYPE: Nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 35 (ii) MOLECULE TYPE: DNA (xi) SÉQUENCE DESCRIPTION: SEQ ID NO:18: ATG TCA TCA GGT GTT TAT GAA TCG ATC ATT CAG ACA AAG 39 GCT TCA GIT TGG GGA TCG ACA GCA TCT GGT AAG TCC ATC 78





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(I) SEQUENCE CHARACTERISTICS: (A) LENGTH: 777 base pairs (B) TYPE: Nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: lihear (II) MOLEGULE TYPE: DNA (XI) SEQUÊNCE DESCRIPTION: SEQ ID NO:19: ATG TOT AAG GTC AAG CTC ACA AAA GAA AAC ATT GTC TCT CIT TTA 45 ÁCT CẦA TỚT GÓA GẮƯ GỮT GAG TÍT GAA GAA GÁC CÁG ÁAC CAG GIC , 90 GCA TTC AAC TTT AAG ACT TIC TGT CAG GAA AAT CIT GAC CIG ATT 135 AAG AAA ATG AĞT ATC ACT TCA TGT TTG ACT TTC TTG AAG AAT CGC 180 CAA GGC ATT ATG AAA GIT GTG AAC CAA AGT GAT TIT ACT TIT GGT 225 ÀAG GTC ACG ATA AAG ÀAA AAT TCT GAA AGA GIT GGÁ GCT AAG GAT 270 ATG ĂCT TIC AGG AGG CTT GÁT AGC ATG ATA AGA GIG AAG CIC ATA 315 GAÁ GÁG ACT GCÁ ÁAC ÁÁT GAG AAT CIT GCT ATT ÁTC AAA GCA AAA 360 ÁTT CCC TCC CAC CCT TTG GTC CAA GCT TAC GGG CTG CCT CTG GCA 405 450 GAT GCA AAA TCT GIG AGA CIT GCT ATA ATG CIT GGA GGT AGT ATC 495 CCT CTC ATT GCT TCT GTT GAC AGC TTC GAG ATG ATC AGT GTT GTT 540 CIT GCC ATA TAT CAA GAT GCA AAG TAC AAG GAG TTA GGG ATT GAA CCA ACT AAG TAC ÁAC ACT AAG GAA GCT CTG GGG AAG GTT TGC ACA 585 630 GTG CTT AAA AGC AAA GGÁ TTT ACA ATG GAT GAT GCA CAG ATA AAC AAA GGG AAA GAA TAT GCC AAG ATA CTT AGT TCT TGC AAT CCC AAT 675 GCT AAG GGA AGC ATT GCT ATG GAC TAT TAC AGT GAT AAT CIT GAC 720 AAA TIC TAT GAA ATG TIT GGA GIT AAG AAA GAG GCC AAG ATT GCT 765 GGT GTT GCA TAA 777 (I) SEQUENCE CHARACTERISTICS: (A) LENGTH: 40 base pairs

(2) INFORMATION FOR SEQ ID NO:20:

(B) TYPE:

Nucleic acid

(C) STRANDEDNESS:

single

(D) TOPOLOGY:

linear

(ii) MOLECULE TYPE:

DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

TACTIATCIA GAACCATGGA CAAAGCAAAG ATTACCAAGG

(2) INFÖRMATION FOR SEQ ID NO:21: 35

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH:

42 base pairs

	(B) TYPE: N (C) STRANDEDNESS: (D) TOPOLOGY: III	ucleic acid single	
	(ii) MOLECULE TYPE: D		
5	(xi) SEQUENCE DESCRIPTION		
3		n. segid no.21. Tat ticaaataat tiataaaag	C AC 42
	(2) ÎNFORMATION FOR SEQ ID NO:		
	(I) SEQUENCE CHARACTER!		
	(A) LENGTH: 3 (B) TYPE: N	6 base pairs	,
10	(B) TYPE: N	lucleic acid	
·	(C) STRANDEDNESS:	single `	
	(D) TOPÖLOGY: II		
	(ii) MÖLECULE TYPE: D		
خ د	(xi) SEQUENCE DESCRIPTIO		
15		TAA CACACTAAGC AAGCAC 3	6
	(2) INFORMATION FOR SEQ ID NO:		
	(i) SEQUENCE CHARACTERI	STICS:	
ì	(A) LENGTH: 4 (B) TYPE: N	6 base pairs	
20	(C) STRANDEDNESS:	UCIEIC ACIO	
	(D) TOPOLOGY: 1	near	
	(ii) MOLECULE TYPE:		
	(xi) SEQUENCÉ DESCRIPTIO	N: SEQ ID NO:23:	
	AGCIAATCIA GAACCATGGA TGACTCA		6
25	(2) INFORMATION FOR SEQ ID NO:		
	(i) SEQUENCE CHARACTERI	STICS:	
	(A) LENGTH: 2	22 base pairs	
	(B) TYPE:	Nucleic acid	
30	(C) STRANDEDNESS:	•	
30		inear	
		ONA	
	(xi) SEQUENCE DESCRIPTION		
•	CCCACIATCC TTCGCAZ		
	(2) INFORMATION FOR SEQ ID NO:		
35	(i) SEQUENCE CHARACTER	ISTICS:	×.
		39 base pairs	`
		Nucleic acid	
	(C) STRÅNDEDNESS: (D) TOPÖLOGY: I		

	(ii) MOLECULE TYPE: [AVA	
	(xí) SEQUENCE DESCRIPTIC	N: SEQ ID NO:25:	
	TACAGIGGĂÎ ČCĂTGG	TTÁÁ GGTAATOCÁT ÁGGCTTGAC 39	
	(2) INFÖRMATION FOR SEQ ID NO	26:	
5		10 bäse pairs Nucleic acid single	
10	(ii) MÓLECULE TYPÉ:	DNA .	
	(xi) SEQUENCE DESCRIPTION	DN: SEQ ID NO:26:	
	AGCTÁACCAT GGTTAA	GCIC ACIAAGGAAA GCATTGITGC 40	
	(2) INFORMATION FOR SEQ ID NO	:27:	
15	(i) SEQUENCE CHARACTER (A) LENGTH: (B) TYPE: (C) STRANDEDNESS	46 base pairs Nucleic acid : single	
$\frac{\partial}{\partial x}$	(D) TOPOLOGY:		
20	(II) MÓLECULÉ TÝPE:		
20	(xi) SEQUENCE DESCRIPTI	on. seg id no.27. Igacicacia aggaaagcai igiigc 4	c
	(2) INFÖRMÁTIÓN FOR SEQ ID NO		O
25	(i) SEQUÊNCE CHARACTEI (A) LENGTH:	RISTICS: 36 base pairs Nucleic acid : single linear	
	(xi) SEQUENCE DESCRIPT		
30	• •	GITAA CACACIAAGC AAGCAC 36	
30	(2) INFORMATION FOR SEQ ID N		
35	(i) SEQUENCE CHARACTE (A) LENGTH: (B) TYPE: (C) STRANDEDNES: (D) TOPOLOGY:	RISTICS: 39 base pairs Nucleic acid 5: single	`
	(ii) MOLECULE TYPE:	DNA	
	(xi) SEQUENCE DESCRIPT	ION: SEQ ID NO:29:	
	TACAGITCIA GAACO	ATGGA TGATGCAAAG TCTGTGAGG 39	
	•		

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(2) INFORMATION FOR SEQ ID NO:30:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH:

49 base pairs

(B) TYPE:

(C) STRANDEDNESS:

Nucleic acid

(D) TOPOLOGY:

linear

(II) MÓLECULE TYPE:

DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

AGATICICIA GACCATGGIG ACTIGATGAG CAAAGICIGI GAGGCIIGC 49

Thus while we have illustrated and described the preferred embodiments of our invention, it is to be understood that this invention is capable of variation and modification, and we therefore do not wish to be limited to the precise terms set forth, but desire to avail ourselves of such changes and alterations which may be made for

- adapting the invention to various usages and conditions. Such variations and modifications, for example, would include the substitution of structurally similar nucleic acid sequences in which the difference between the sequence shown and the variation sequence is such that little if any advantages are available with the variation
- sequence, i.e. that the sequences produce substantially similar results as described above. Thus, changes in sequence by the substitution, deletion, insertion or addition of nucleotides (in the nucleotide sequences) or amino acids (in the peptide sequences) which do not substantially alter the function of those sequences specifically
- described above are deemed to be within the scope of the present invention. In addition, it is our intention that the present invention may be modified to join the N genes of various isolates that provide resistance or immunity to *Tospovirus* infection of plants according to the present invention into a single cassette, and to use this cassette as a transgene in order to provide broad resistance to the Tospoviruses.
 - a transgene in order to provide broad resistance to the Tospoviruses, especially to TSWV-BL, TSWV-B, and INSV. Accordingly, such changes and alterations are properly intended to be within the full range of equivalents, and therefore within the purview of the following claims.

Having thus described our invention and the manner and a process of making and using it in such full, clear, concise and exact terms so as

to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same;

j,

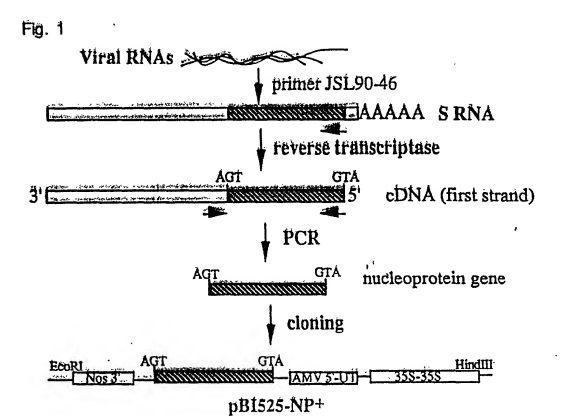
We Claim:

1. An isolated nucleotide sequence which is selected from the

Travera	
group	=-
AAATTGTCTT GCAGTGAAAT CTCTGCTCAT GTTAGCAGAA AACAACATCA	50
TECCTAACIC TCAACCITTT GICAAACCIT CIACIGATIC TAATITCAAG	100
CIGAGCEICT GECTAAGGET TCCAAAGGET TIGAAGCAGA TITICCATTCA	150
GAAATTGTTC AAGGTTGGAG GAGATGAAAC AAATAAAACA TTTTATTTAT	200
CIATTGCCIG CATTCCAAAC CATAACAGIG TIGAGACAGC TITTAAACATT	250
ACIGITATTT GCAACCATCA GCTCCCAATT CGTAAATGTA AAACTCCTTT	300
TCAATTATCA ATGATCITTT CIGATTTAAA GGAGCCTTAC AACATTATTC	350 ′
ATCATECTIC ATATOCOCAA AGGATTGTTC ATGCTCTGCT TGAAACTCAC	400
ACATCTTITE CACAACTTCT TIECAACAAC TIECAACAAG ATGIGATCAT	450
CTAGASCITG AACAACCATG AGCTAACTCC TGGAAAGTTA GATTTAGGTG	500
AAATAACTTT GAATTACAAT GAAGACGCCT ACAAAAGGAA ATATTTCCTT	550
TCAAAAACAC TTCAATCTT TCCATCTAAC ATACAAACTA TGTCTTATTT	600
AGACAGCATC CAAATCCCTT CCTGGAAGAT AGACTTTGCC AGGGGAGAAA	650
TTÄÄÄÄTTTC TCCACÄÄTCT ÄTTTCAGTTG CAAAATCTTT GTTAAATCTT	700
GAÍTTÍAAGCG GÉATTIAAAA GAAAGAATCT AAGATTAAGG AAGCATATGC	750
TICAGGATCA AAATGATCTT GCTGTGTCCA GCTTTTTCTA ATTATGTTAT	800
GITTATTIC TITCITTACT TATAATIATT TITCIGITIG TCATTICTT	850
CAAATTCCTC CTGTCTAGTA GAAACCATAA AAACAAAAAT AAAAATAAAA	900
TAÄÄÄÍTCAAA ÄTÄÄÄÄÍTÄÄÄ ÄÄTCÄÄÄAAA TGAAATÄAAA GCAACAAAÁA	950
AATTAAAAA CAAAAAACCA AAAAAGATCC CGAAAGGACA ATTTTGGCCA	1000
is a constitution of the c	1050
Let be All Company that the second of the se	1100
CONTRACTOR CONTRACTOR AND AND AND AND AND AND AND AND AND AND	1150
TAAÂĆÁCÁCT TÁŤTTÁÁÁÁŤ ŤŤÁÅCÁCACT AAGCAAGCACÁ AACAATAAA	1200
GATAAAGAAA GCITTATTATA TTTTATAGGCT TTTTTTATAAT TTAACTTACA	
GCTGCTTTA AGCAAGTTCT GTGAGTTTTG CCTGTTTTTT AACCCCAAAC	
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ATTATIGGCAA GCCTCACAGA CTTTGCATCA TCAAGAGGTA ATCCATAGGC	
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CAGAATICC 1709;	1,00
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TITATAGGCT TITTTATAAT TTAACTTACA GCIGCTTTTA AGCAAGTTCT	
GTGAGITITG CCIGITITIT AACCCCAAAC ATTICATAGA ACTIGITAAG	
GGITTCACIG TAATGITCCA TAGCAATACT TCCTTTAGCA TTAGGATTGC	200
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TICATTICAA ATGCTTTTCT TTTCAGCACA GIGCAAACIT TTCCTAAGGC	
TICCIGGIG TCATACITCT TIGGGTOGAT CCCGAGATCC TIGIATTITG	
**************************************	350

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and AGAGCAATTIG GGICATTITIT TATTICTAAAT CGAACCTCAA CTAGCAAATC 50 TCÁGÁÁCÍGÍ AÁTAÁGCÁCÁ ÁGAGCÁCAÁG ÁGCCÁCÁATG TCATCAGGTG 100 TTTATCAATC GATCATTCAG ACAAAGGCTT CAGTTTGGGG ATCGACAGCA 150 TCTGGTAAGT CCATCGTGGA TTCTTACTGG ATTTATGAGT TTCCAACTGG 200 TÜĞÜĞACİĞ ĞİTCAAACİC AĞİTİĞFACIC TGATTICGAGG AĞCAAAAĞTA 250 GCTTCGGCTA CACTTCAAAA ATTGGTGATA TICCTGCTGT AGAGGAGGAA 300 ATTITATION AGAAGITICA TATOCCAGIG TITGATGATA TIGATTICAG 350 CATCAATATC AATGATTCTT TCTTGGCAAT TTCTGTTTGT TCCAACACAG 400 TTÄACACCAA TGGAGTGAAG CATCAGGGTC ATCTTAAAGT TCTTTCTCTT 450 GOCCAATTICC ATCCCTTIGA ACCTGTGATG AGCAGGTCAG AGATTGCTAG 500 CAGATTOCGG CTCCAAGAAG AAGATATAAT TOCTGATGAC AAATATATAT 550 CIGCIGCIAA CAAGGGATCT CICTCCTGTG TCAAAGAACA TACTTACAAA 600 GTÖGAAÁTGÁ GCCACÁÁTCÁ GGCTTTAGGC ÁAAGTGAATG TTCTTTCTCC 650 TAÁCÁGAAÁT GTTCÁTGAGT GGCTGTATAG TITCÁAACCA AATTTCÁACC 700 AGATOGÁAÁG TAATAÁCÁGA ÁCIGTAAATT CICTTGCÁGT CÁAATCITTG 750 CÍCATGCTA CAGAAAACAA CÁTTATGCCT AACTCTCAAG CTTTTGTTAA 800 AGCITCIACT GATTICICATT TTAAGTTGAG CCTTTGGCTG AGAATTCCAA 850 AAGTTTTGÁÁ GCÁÁÁTÁGCC ÁTÁCAGAGC TCTTCÁAGTT TGCÁGGÁGAC 900 GAÀÀCCGGTA ÀÀAGTTTCTÁ TTTGTCTATT GCATGCATCC CAAATCACAA 950 CAGTGTGGAA ACAGCTTTAA ATGTCACTGT TATATGTAGA CATCAGCTTC 1000 CAATOCCTAA GICCAAAGCT CCTTTTGAAT TATCAATGAT TTTCTCCGAT 1050 CIGAAAGAGC CTTACAACAC TGTGCATGAT CCTTCATATC CTCAAAGGAT 1100 TGTTCATGCT TTGCTTGAGA CTCACACTTC CITTGCACAA GTTCTCTGCA 1150 ACAAGCIGCA ÁGAAGAIGIG ÁTCATATATA CIATAAACAG CCCIGAACIA 1200 ACCCCÁGCIA ÁGCTGGÁTICT ÁGGTGAAAGA ACCTTGAACT ACAGTGAAGA 1250 TGCTTCGAAG AAGAAGTATT TTCTTTCAAA AACACTCGAA TGCTTGCCAG 1300 TAAATGIGCA GACTATGICT TATTIGGATA GCATCCAGAT TCCTTCATGG 1350 AAGATAGACT TTGCCAGAGG AGAGATCAGA ATCTCCCCTC AATCTACTCC 1400 TATTGCAAGA TCTTTGCTCA AGCTGGATTT GAGCAAGATC AAGGAAAAGA 1450 AGICCITIGAC THEGGAAACA TOCAGCTATE ATCTAGAATA AAAGTEGCTC 1500 ATACTACICT AAGTAGTATT TGTCAACTTG CTTATCCTTT ATGTTGTTTA 1550 TITCITTAA ATCHAAAGIA AGITAGATIC AAGIAGITTA GTATGCTATA 1600 GCATTATTAC AAAAAATACA AAAAATACA AAAAATATAA 1650 AÁÁACCCÁAÁ AÁGATCCCAÁ ÁAGGGÁCGAT TIGGITGATT TACICTGITT 1700 TAGGCTTATC TAAGCTGCTT TIGTTTGAGC AAAATAACAT TGTAACATGC 1750 AATAACIGGA ÀTTTAÁAGIC CTÁAAAGAG TITCAAAGGA CAGCTTAGCC 1800 AAATTIGGIT TITGITTITG TITTITTIGIT TITTIGITTIT TIGITTATT 1850 1900 THATHITA GILLATITE TGILLIGIT ATTITATT TEATTTATT THOTTHATT THATHTATAT ATATATCAAA CACAATCCAC ACAATAATT 1950 THANTITICAA ACAITICÍACT GATTTAACAC ACITAGOCTG ACITTATCAC 2000 2050 2100 CTTAGTATTA TGCATCTCTT AATTTAACACA CTITTAATAAT ATGCATCTCT GAATCAGOCT TAAAGAAGCT TTTATGCAAC AOCAGCAATC TTGGCCTCTT 2150 TCITAACIOC AAACATTICA TAGAATTIGT CAAGATTATC ACIGIAATAG 2200 TOTATAGCAA TOCTTCCCTT AGCATTGGGA TIGCAAGAAC TAAGIATCIT 2250 2300 GOCATATICE TECCCETTICE TEATCEGIGC ATCATCCATE GEARATCCET TECTTITAAG CACTETECAA ACCITICCOCA GAGCITCCIT AGIGITGIAC 2350 THAGITGGIT CAATOCCIAA CICCITGIAC TITICCATOIT GATATATGGO 2400 AAGAACAACÁ CTGÁTCÁTCT CGAAGCTGTC AACÁGAAGCA ATGAGAGGGA 2450 TACTACCICC AAGCATTATA GCAAGICICA CAGATTTIGC ATCIGCCAGA 2500 2550 GGCÄGCCCGT ÀAGCTTGGAC CÂAAGGGTGG GAGGCAATTT TIGCTTTGAT 2600 AATAGCAAGA TICTCATIGT TIGCAGICIC TICIATGAGC TICACICITA TCATGCTATC AAGCCTCCTG AAAGTCATAT CCTTAGCTCC AACTCTTTCA 2650 GAATTITICT TTATOGIGAC CITACCAAAA GIAAAATCAC TTIGGITCAC 2700 2750 AACTITICATA ATGCCTTGGC GATTCTTCAA GAAAGTCAAA CATGAAGTGA TACTCATTIT CTTAATCAGG TCAAGATTIT CCTGACAGAA AGTCTTAAAG 2800 TIGAATGCGA CCIGGTICTG GICTTCTTCA AACTCAACAT CIGCAGATTG 2850 2900 AGITAÁAÁGA GAGACAÁTGT TTTCTTTTGT GAGCTTGACC TTAGACATGG TGGCAGITHA GATCHAGACC TITICTOGAGA GATAAGATTC AAGGTGAGAA 2950 AGIGCAACAC TGIAGACCGC GGICGITACT TATCCIGITA ATGIGATGAT 3000 TIGIATIGCT GAGIATTAGG TITTTIGAATA AAATTGACAC AATTGCTCT 3049

- 2. A plant susceptible to infection by *Tospoviruses* which has a transgene inserted into its genome to render it resistant to infection by Tospoviruses, said transgene being selected from the group consisting of the nucleoprotein gene of TSWV-BL, TSWV-10W, INSV-LI, TSWV-B, a *Tospovirus*, said transgene consisting of partial or full length nucleoprotein gene sequences from TSWV-BL, TSWV-10W, TSWV-B, INSV-Beg and INSV-IL, the translatable or untranslatable sequences of said nucleoprotein gene sequences, and the sense or antisense sequences of said nucleoprotein gene sequences.
- 3. A method for providing a host plant with resistance to infection by *Tospoviruses* which comprises insetting a transgene into the host plant which gene is selected from the nucleoprotein gene of TSWV-BL, TSWV-10W, INSV-Beg, INSV-LI, TSWV-B, or mixtures of nucleotide sequences taken from the nucleoprotein gene.



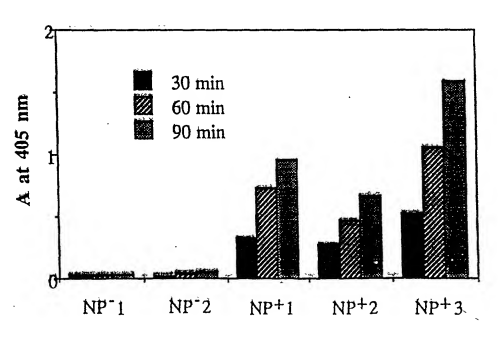


Fig. 2

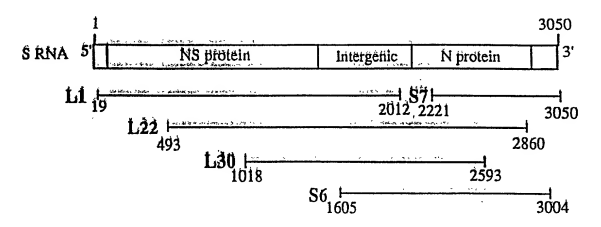
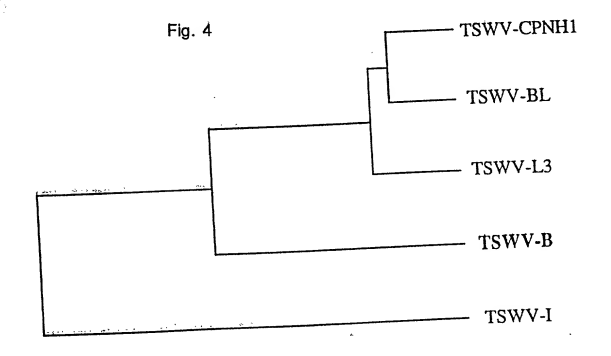
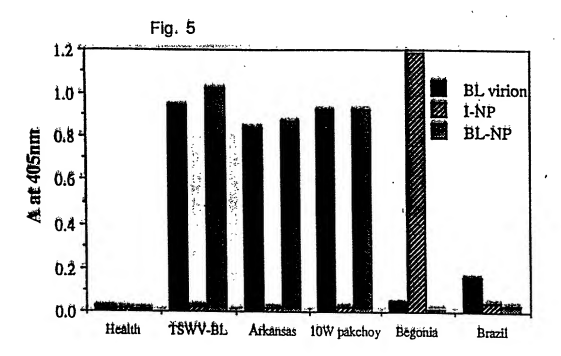
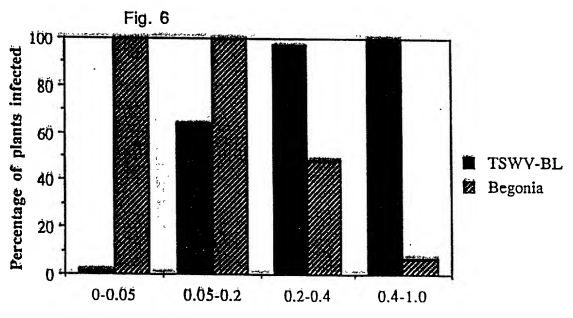


Fig. 3



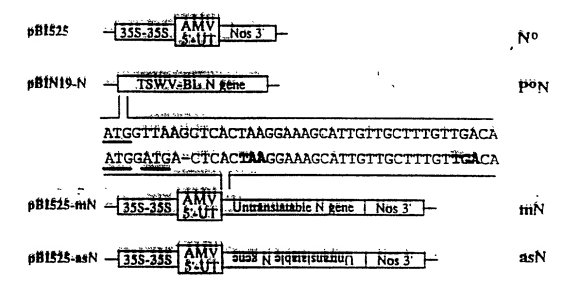




The NP accumulated in plants (ELISA, A at 405nm)

题、图文、 1780 · 18 (图) 图

Fig. 7



		Plant line
	ATG TSW.VBL N gene TAA	
pbini9-in	- 1970 Annual -	1N
	<u>ATG</u> GTTAAGGTCACTAAGGAAAGCATTGTTGCTTTGTTGACA	
	ÀTGGÀTGÀ CTCÀ TÀ ÀGGÀA ÀGC ÀTTGTTGCTTTGTGACA	•
pBini9-in'	committee on whitehearter - and stronger - TAA	1N'
pBIN19-1N	AAT OTA	1N -
pBin19-2n	TAA	2N
	ATGGATGATGCAAAGTCTGTGAGGCTTGCCATAATG	
	ATGGTGACTTGATGAGCAAAGTCTGGAGGCTTGCCATAATG	
pBIN19-2N'	TM	2N'
pbini9-2n	AAT TTA	2N -

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ATTORNEY'S DOCKET NUMBER COMBINED DECLARATION FOR PATENT

I (II	APPLICATION AND POWER OF ATTORNEY cludes Reference to PCT International Applications)	19603/10301
As a be	low named inventor, I hereby declare that:	
My resi	dence, post office address and citizenship are as stated	below next to my name.
or an o	ve I am the original, first and sole inventor (if only original, first and joint inventor (if plural names are l subject matter which is claimed and for which a patent id: TOMATO SPOTTED WILT VIRUS	isted below)
the spe	cification of which (check only one item below):	
[]	is attached hereto.	
[]	was filed as United States application Serial No.	
	onand was amended	(1.5
	and was amended on	(if applicable).
(x)	and was amended on was filed as PCT international application	(if applicable).
[x]	and was amended on was filed as PCT international application NumberPCT/US94/01046 on _January 27, 1994	(if applicable).
[x]	and was amended on was filed as PCT international application NumberPCT/US94/01046	(if applicable).
I hereb	and was amended on was filed as PCT international application NumberPCT/US94/01046 on _January 27, 1994	of the above-identified
I hereb	and was amended on	of the above-identified int referred to above.

PRIOR FOREIGN/PCT APPLICAT	TION(S) AND ANY PRIOR	ITY CLAIMS UNDER 3	5 U.S.C. 119:
COUNTRY (IF PCT, indicate "PCT")	APPLICATION NUMBER	DATE OF FILING (day, month, year)	PRIORITY CLAIMED UNDER 35 USC 119
			[] YES [] NO
			[] YES [] NO
		I	PAGE 1 OF 2

COMBINED DECLARATION FOR PATENT
APPLICATION AND POWER OF ATTORNEY (Continued)
(Includes Reference to PCT International Applications)

ATTORNEY'S DOCKET NUMBER 19603/10301

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) or PCT international application(s) designating the United States of America that is/are listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in that/those prior application(s) in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, § 1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT International filing date of this application:

PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS DESIGNATING THE U.S. FOR BENEFIT UNDER 35 U.S.C. 120:

U.S. APPLICATIONS				STATUS (Che	ck One)	
U.S. APPLICATION NUMBER		\v.s	. FILING DATE	PATENTED	PENDING	ABANDONED
08/010,410		29 January 1993			· x	
PCT APPL	ICATIONS DESIGNATI	NG THE	U.S.			
PCT APPLICATION NO.	PCT FILING DATE		SERIAL NUMBERS			
PCT/US94/01046	27 January 1994					

DOWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number)

Michael L. Goldman, Reg. No. 30,727; Susan J. Timian, Reg. No. 34,103

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	FULL NAME OF INVENTOR	FAMILY NAME Gonsalves	FIRST GIVEN NAME Dennis	SECOND GIVEN NAME	
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	POST OFFICE ADDRESS	P.O. ADDRESS 595 Castle Street	CITY Geneva	STATE & ZIP CODE/COUNTRY New York 14456	
	FULL NAME OF INVENTOR	FAMILY NAME Pang	FIRST GIVEN NAME Sheng-Zhi	SECOND GIVEN NAME	
2 0 2	RESIDENCE & CITIZENSHIP	CITY - Chesterfield	STATE/FOREIGN COUNTRY 5.8	COUNTRY OF CITIZENSHIP China	
	POST OFFICE ADDRESS	P.O. ADDRESS 5.7 666 West North Street 893 J Foxsprings Dr.	CITY sp Ganova Chesterfield	STATE & ZIP 5 ? . CODE/COUNTRY M.55 puri New York 14456 63148	

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section IOO1 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

SIGNATURE OF INVENTOR 201	SIGNATURE OF INVENTOR 2025	
PATE September 14, 1985	DATE Sept. 19, 1995	

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: Gonsalves, Dennis Pang, Sheng-Zhi
 - (ii) TITLE OF INVENTION: TOMATO SPOTTED WILT VIRUS
 - (iii) NUMBER OF SEQUENCES: 30
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Nixon Peabody LLP
 - (B) STREET: Clinton Square, P.O. Box 1051
 - (C) CITY: Rochester
 - (D) STATE: New York
 - (E) COUNTRY: U.S.A.
 - (F) ZIP: 14603
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/495,484
 - (B) FILING DATE: 27-JAN-1994
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Goldman, Michael L.
 - (B) REGISTRATION NUMBER: 30,727
 - (C) REFERENCE/DOCKET NUMBER: 19603/10303
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (716) 263-1304
 - (B) TELEFAX: (716) 263-1600
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 25 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

(2)	INFORMATION FOR SEQ ID NO:2:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
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(2)	INFORMATION FOR SEQ ID NO:3:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 32 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
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(2)	INFORMATION FOR SEQ ID NO:4:	
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	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
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(2)	INFORMATION FOR SEQ ID NO:5:	
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(A) LENGTH: 2216 base pairs (B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

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AACAAAAAAT	TAAAAAACGA	AAAACCAAAA	AGACCCGAAA	GGGACCAATT	TGGCCAAATT	1020
TGGGTTTTGT	TTTTGTTTTT	TGTTTTTTGT	TTTTTATTTT	TTATTTTATT	TTTATTTTAT	1080
TTTATTTTTA	TTTTATTTT	ATTTTATTTA	TTTTTTGTTT	TCGTTGTTTT	TGTTATTTTA	1140
TTATTTATTA	AGCACAACAC	ACAGAAAGCA	AACTTTAATT	AAACACACTT	ATTTAAAATT	1200
TAACACACTA	AGCAAGCACA	AGCAATAAAG	ATAAAGAAAG	CTTTATATAT	TTATAGGCTT	1260
TTTTATAATT	TAACTTACAG	CTGCTTTCAA	GCAAGTTCTG	CGAGTTTTGC	CTGCTTTTTA	1320
ACCCCGAACA	TTTCATAGAA	CTTGTTAAGA	GTTTCACTGT	AATGTTCCAT	AGCAACACTC	1380
CCTTTAGCAT	TAGGATTGCT	GGAGCTAAGT	ATAGCAGCAT	ACTCTTTCCC	CTTCTTCACC	1440

TGATCTTCAT	TCATTTCAAA	TGCTTTGCTT	TTCAGCACAG	TGCAAACTTT	TCCTAAGGCT	1500
TCCTTGGTGT	CATACTTCTT	TGGGTCGATC	CCGAGGTCCT	TGTATTTTGC	ATCCTGATAT	1560
ATAGCCAAGA	CAACACTGAT	CATCTCAAAG	CTATCAACTG	AAGCAATAAG	AGGTAAGCTA	1620
CCTCCCAGCA	TTATGGCAAG	TCTCACAGAC	TTTGCATCAT	CGAGAGGTAA	TCCATAGGCT	1680
TGAATCAAAG	GATGGGAAGC	AATCTTAGAT	TTGATAGTAT	TGAGATTCTC	AGAATTCCCA	1740
GTTTCTTCAA	CAAGCCTGAC	CCTGATCAAG	CTATCAAGCC	TTCTGAAGGT	CATGTCAGTG	1800
CCTCCAATCC	TGTCTGAAGT	TTTCTTTATG	GTAATTTTAC	CAAAAGTAAA	ATCGCTTTGC	1860
TTAATAACCT	TCATTATGCT	CTGACGATTC	TTTAGGAATG	TCAGACATGA	AATAACGCTC	1920
ATCTTCTTGA	TCTGGTCGAT	GTTTTCCAGA	CAAAAAGTCT	TGAAGTTGAA	TGCTACCAGA	1980
TTCTGATCTT	CCTCAAACTC	AAGGTCTTTG	CCTTGTGTCA	ACAAAGCAAC	AATGCTTTCC	2040
TTAGTGAGCT	TAACCTTAGA	CATGATGATC	GTAAAAGTTG	TTATATGCTT	TGACCGTATG	2100
TAACTCAAGG	TGCGAAAGTG	CAACTCTGTA	TCCCGCAGTC	GTTTCTTAGG	TTCTTAATGT	2160
GATGATTTGT	AAGACTGAGT	GTTAAGGTAT	GAACACAAAA	TTGACACGAT	TGCTCT	2216

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1709 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

AAATTCTCTT	GCAGTGAAAT	CTCTGCTCAT	GTTAGCAGAA	AACAACATCA	TGCCTAACTC	60
TCAAGCTTTT	GTCAAAGCTT	CTACTGATTC	TAATTTCAAG	CTGAGCCTCT	GGCTAAGGGT	120
TCCAAAGGTT	TTGAAGCAGA	TTTCCATTCA	GAAATTGTTC	AAGGTTGCAG	GAGATGAAAC	180
AAATAAAACA	TTTTATTTAT	CTATTGCCTG	CATTCCAAAC	CATAACAGTG	TTGAGACAGC	240
TTTAAACATT	ACTGTTATTT	GCAAGCATCA	GCTCCCAATT	CGTAAATGTA	AAACTCCTTT	300
TGAATTATCA	ATGATGTTTT	CTGATTTAAA	GGAGCCTTAC	AACATTATTC	ATGATCCTTC	360
ATATCCCCAA	AGGATTGTTC	ATGCTCTGCT	TGAAACTCAC	ACATCTTTTG	CACAAGTTCT	420

T'	IGCAACAAC	TTGCAAGAAG	ATGTGATCAT	CTACACCTTG	AACAACCATG	AGCTAACTCC	480
T(GGAAAGTTA	GATTTAGGTG	AAATAACTTT	GAATTACAAT	GAAGACGCCT	ACAAAAGGAA	540
A.	PATTTCCTT	TCAAAAACAC	TTGAATGTCT	TCCATCTAAC	ATACAAACTA	TGTCTTATTT	600
A	GACAGCATC	CAAATCCCTT	CCTGGAAGAT	AGACTTTGCC	AGGGGAGAAA	TTAAAATTTC	660
T	CCACAATCT	ATTTCAGTTG	CAAAATCTTT	GTTAAATCTT	GATTTAAGCG	GGATTAAAAA	720
G	AAGAATCT	AAGATTAAGG	AAGCATATGC	TTCAGGATCA	AAATGATCTT	GCTGTGTCCA	780
G	CTTTTTCTA	ATTATGTTAT	GTTTATTTTC	TTTCTTTACT	TATAATTATT	TTTCTGTTTG	840
TO	CATTTCTTT	CAAATTCCTC	CTGTCTAGTA	GAAACCATAA	AAACAAAAAT	AAAAATAAAA	900
ΤZ	AAATCAAA	ATAAAATAAA	AATCAAAAAA	TGAAATAAAA	GCAACAAAA	AATTAAAAAA	960
CZ	AAAAACCA	AAAAAGATCC	CGAAAGGACA	ATTTTGGCCA	AATTTGGGGT	TTGTTTTTGT	1020
T	TTTTGTTTT	TTTGTTTTTT	GTTTTTATTT	TTATTTTTAT	TTTTATTTTT	ATTTTATTTT	1080
A7	TTTATGTT	TTTGTTGTTT	TTGTTATTTT	GTTATTTATT	AAGCACAACA	CACAGAAAGC	1140
ΑZ	ACTTTAAT	TAAACACACT	TATTTAAAAT	TTAACACACT	AAGCAAGCAC	AAACAATAAA	1200
GÆ	ATAAAGAAA	GCTTTATATA	TTTATAGGCT	TTTTTATAAT	TTAACTTACA	GCTGCTTTTA	1260
ΑC	SCAAGTTCT	GTGAGTTTTG	CCTGTTTTTT	AACCCCAAAC	ATTTCATAGA	ACTTGTTAAG	1320
GG	TTTCACTG	TAATGTTCCA	TAGCAATACT	TCCTTTAGCA	TTAGGATTGC	TGGAGCTAAG	1380
TA	TAGCAGCA	TACTCTTTCC	CCTTCTTCAC	CTGATCTTCA	TTCATTTCAA	ATGCTTTTCT	1440
ТТ	TCAGCACA	GTGCAAACTT	TTCCTAAGGC	TTCCCTGGTG	TCATACTTCT	TTGGGTCGAT	1500
CC	CCGAGATCC	TTGTATTTTG	CATCCTGATA	TATAGCCAAG	ACAACACTGA	TCATCTCAAA	1560
GC	CTATCAACT	GAAGCAATAA	GAGGTAAGCT	ACCTCCCAGC	ATTATGGCAA	GCCTCACAGA	1620
CI	TTGCATCA	TCAAGAGGTA	ATCCATAGGC	TTGAATCAAA	GGGTGGGAAG	CAATCTTAGA	1680
тт	TGATAGTA	TTGAGATTCT	CAGAATTCC				1709

(2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 260 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Gln Val Glu Ser Asn Asn Arg Thr Val Asn Ser Leu Ala Val Lys Ser

1 5 10 15

Leu Leu Met Ser Ala Glu Asn Asn Ile Met Pro Asn Ser Gln Ala Ser 20 25 30

Thr Asp Ser His Phe Lys Leu Ser Leu Trp Leu Arg Val Pro Lys Val 35 40 45

Leu Lys Gln Val Ser Ile Gln Lys Leu Phe Lys Val Ala Gly Asp Glu 50 55 60

Thr Asn Lys Thr Phe Tyr Leu Ser Ile Ala Cys Ile Pro Asn His Asn 65 70 75 80

Ser Val Glu Thr Ala Leu Asn Ile Thr Val Ile Cys Lys His Gln Leu 85 90 95

Pro Ile Arg Lys Cys Lys Ala Pro Phe Glu Leu Ser Met Met Phe Ser 100 105 110

Asp Leu Lys Glu Pro Tyr Asn Ile Val His Asp Pro Ser Tyr Pro Lys
115 120 125

Gly Ser Val Pro Met Leu Trp Leu Glu Thr His Thr Ser Leu His Lys 130 135 140

Asn Leu Glu Leu Thr Pro Gly Lys Leu Asp Leu Gly Glu Arg Thr Leu 165 170 175

Asn Tyr Ser Glu Asp Ala Tyr Lys Arg Lys Tyr Phe Leu Ser Lys Thr 180 185 190

Leu Glu Cys Leu Pro Ser Asn Thr Gln Thr Met Ser Tyr Leu Asp Ser 195 200 205

Ile Gln Ile Pro Ser Trp Lys Ile Asp Phe Ala Arg Gly Glu Ile Lys 210 215 220

Ile Ser Pro Gln Ser Ile Ser Val Ala Lys Ser Leu Leu Lys Leu Asp 225 230 235 240

Leu Ser Gly Ile Lys Lys Lys Glu Ser Lys Val Lys Glu Ala Tyr Ala 245 250 255

Ser Gly Ser Lys 260

(2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 858 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

	TTAACACACT	AAGCAAGCAC	AAACAATAAA	GATAAAGAAA	GCTTTATATA	TTTATAGGCT	60
	TAATATTTT	TTAACTTACA	GCTGCTTTTA	AGCAAGTTCT	GTGAGTTTTG	CCTGTTTTTT	120
	AACCCCAAAC	ATTTCATAGA	ACTTGTTAAG	GGTTTCACTG	TAATGTTCCA	TAGCAATACT	180
,	ICCTTTAGCA	TTAGGATTGC	TGGAGCTAAG	TATAGCAGCA	TACTCTTTCC	CCTTCTTCAC	240
4	CTGATCTTCA	TTCATTTCAA	ATGCTTTTCT	TTTCAGCACA	GTGCAAACTT	TTCCTAAGGC	300
	PTCCCTGGTG	TCATACTTCT	TTGGGTCGAT	CCCGAGATCC	TTGTATTTTG	CATCCTGATA	360
,	TATAGCCAAG	ACAACACTGA	TCATCTCAAA	GCTATCAACT	GAAGCAATAA	GAGGTAAGCT	420
i	ACCTCCCAGC	ATTATGGCAA	GCCTCACAGA	CTTTGCATCA	TCAAGAGGTA	ATCCATAGGC	480
	ITGACTCAAA	GGGTGGGAAG	CAATCTTAGA	TTTGATAGTA	TTGAGATTCT	CAGAATTCCC	540
Ž	AGTTTCCTCA	ACAAGCCTGA	CCCTGATCAA	GCTATCAAGC	CTTCTGAAGG	TCATGTCAGT	600
(GCTCCAATC	CTGTCTGAAG	TTTTCTTTAT	GGTAATTTTA	CCAAAAGTAA	AATCGCTTTG	660
(CTTAATAACC	TTCATTATGC	TCTGACGATT	CTTCAGGAAT	GTCAGACATG	AAATAATGCT	720
(CATCTTTTTG	ATCTGGTCAA	GGTTTTCCAG	ACAAAAAGTC	TTGAAGTTGA	ATGCTACCAG	780
Ž	ATTCTGATCT	TCCTCAAACT	CAAGGTCTTT	GCCTTGTGTC	AACAAAGCAA	CAATGCTTTC	840
(CTTAGTGAGC	TTAACCAT					858

(2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2028 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

AAATTCTCTT	GCAGTGAAAT	CTCTGCTCAT	GTTAGCAGAA	AACAACATCA	TGCCTAACTC	60
TCAAGCTTTT	GTCAAAGCTT	CTACTGATTC	TAATTTCAAG	CTGAGCCTCT	GGCTAAGGGT	120
TCCAAAGGTT	TTGAAGCAGA	TTTCCATTCA	GAAATTGTTC	AAGGTTGCAG	GAGATGAAAC	180
AAATAAAACA	TTTTATTTAT	CTATTGCCTG	CATTCCAAAC	CATAACAGTG	TTGAGACAGC	240
TTTAAACATT	ACTGTTATTT	GCAAGCATCA	GCTCCCAATT	CGTAAATGTA	AAACTCCTTT	300
TGAATTATCA	ATGATGTTTT	CTGATTTAAA	GGAGCCTTAC	AACATTATTC	ATGATCCTTC	360
ATATCCCCAA	AGGATTGTTC	ATGCTCTGCT	TGAAACTCAC	ACATCTTTTG	CACAAGTTCT	420
TTGCAACAAC	TTGCAAGAAG	ATGTGATCAT	CTACACCTTG	AACAACCATG	AGCTAACTCC	480
TGGAAAGTTA	GATTTAGGTG	AAATAACTTT	GAATTACAAT	GAAGACGCCT	ACAAAAGGAA	540
ATATTTCCTT	TCAAAAACAC	TTGAATGTCT	TCCATCTAAC	ATACAAACTA	TGTCTTATTT	600
AGACAGCATC	CAAATCCCTT	CCTGGAAGAT	AGACTTTGCC	AGGGGAGAAA	TTAAAATTTC	660
TCCACAATCT	ATTTCAGTTG	CAAAATCTTT	GTTAAATCTT	GATTTAAGCG	GGATTAAAAA	720
GAAAGAATCT	AAGATTAAGG	AAGCATATGC	TTCAGGATCA	AAATGATCTT	GCTGTGTCCA	780
GCTTTTTCTA	ATTATGTTAT	GTTTATTTTC	TTTCTTTACT	TATAATTATT	TTTCTGTTTG	840
TCATTTCTTT	CAAATTCCTC	CTGTCTAGTA	GAAACCATAA	AAACAAAAAT	ААААТАААА	900
TAAAATCAAA	ATAAAATAAA	AATCAAAAAA	TGAAATAAAA	GCAACAAAA	AATTAAAAAA	960
CAAAAAACCA	AAAAAGATCC	CGAAAGGACA	ATTTTGGCCA	AATTTGGGGT	TTGTTTTTGT	1020
TTTTTGTTTT	TTTGTTTTTT	GTTTTTATTT	TTATTTTTAT	TTTTATTTT	ATTTTATTTT	1080
ATTTTATGTT	TTTGTTGTTT	TTGTTATTTT	GTTATTTATT	AAGCACAACA	CACAGAAAGC	1140
AAACTTTAAT	TAAACACACT	TATTTAAAAT	TTAACACACT	AAGCAAGCAC	AAACAATAAA	1200
GATAAAGAAA	GCTTTATATA	TTTATAGGCT	TTTTTATAAT	TTAACTTACA	GCTGCTTTTA	1260
AGCAAGTTCT	GTGAGTTTTG	CCTGTTTTTT	AACCCCAAAC	ATTTCATAGA	ACTTGTTAAG	1320
GGTTTCACTG	TAATGTTCCA	TAGCAATACT	TCCTTTAGCA	TTAGGATTGC	TGGAGCTAAG	1380
TATAGCAGCA	TACTCTTTCC	CCTTCTTCAC	CTGATCTTCA	TTCATTTCAA	ATGCTTTTCT	1440
TTTCAGCACA	GTGCAAACTT	TTCCTAAGGC	TTCCCTGGTG	TCATACTTCT	TTGGGTCGAT	1500
CCCGAGATCC	TTGTATTTTG	CATCCTGATA	TATAGCCAAG	ACAACACTGA	TCATCTCAAA	1560
GCTATCAACT	GAAGCAATAA	GAGGTAAGCT	ACCTCCCAGC	ATTATGGCAA	GCCTCACAGA	1620

CTTTGCATCA	TCAAGAGGTA	ATCCATAGGC	TTGACTCAAA	GGGTGGGAAG	CAATCTTAGA	1680
TTTGATAGTA	TTGAGATTCT	CAGAATTCCC	AGTTTCCTCA	ACAAGCCTGA	CCCTGATCAA	1740
GCTATCAAGC	CTTCTGAAGG	TCATGTCAGT	GGCTCCAATC	CTGTCTGAAG	TTTTCTTTAT	1800
GGTAATTTTA	CCAAAAGTAA	AATCGCTTTG	CTTAATAACC	TTCATTATGC	TCTGACGATT	1860
CTTCAGGAAT	GTCAGACATG	AAATAATGCT	CATCTTTTTG	ATCTGGTCAA	GGTTTTCCAG	1920
ACAAAAAGTC	TTGAAGTTGA	ATGCTACCAG	ATTCTGATCT	TCCTCAAACT	CAAGGTCTTT	1980
GCCTTGTGTC	AACAAAGCAA	CAATGCTTTC	CTTAGTGAGC	TTAACCAT		2028

- (2) INFORMATION FOR SEQ ID NO:10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 21 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

TTCTGGTCTT CTTCAAACTC A

21

- (2) INFORMATION FOR SEQ ID NO:11:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

CTGTAGCCAT GAGCAAAG

18

- (2) INFORMATION FOR SEQ ID NO:12:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 467 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Met Ser Ser Gly Val Tyr Glu Ser Ile Ile Gln Thr Lys Ala Ser Val 1 5 10 15

Trp Gly Ser Thr Ala Ser Gly Lys Ser Ile Val Asp Ser Tyr Trp Ile 20 25 30

Tyr Glu Phe Pro Thr Gly Ser Pro Leu Val Gln Thr Gln Leu Tyr Ser 35 40 45

Asp Ser Arg Ser Lys Ser Ser Phe Gly Tyr Thr Ser Lys Ile Gly Asp 50 55 60

Ile Pro Ala Val Glu Glu Glu Ile Leu Ser Gln Asn Val His Ile Pro 65 70 75 80

Val Phe Asp Asp Ile Asp Phe Ser Ile Asn Ile Asn Asp Ser Phe Leu 85 90 95

Gln Gly His Leu Lys Val Leu Ser Leu Ala Gln Leu His Pro Phe Glu 115 120 125

Pro Val Met Ser Arg Ser Glu Ile Ala Ser Arg Phe Arg Leu Gln Glu 130 135 140

Glu Asp Ile Ile Pro Asp Asp Lys Tyr Ile Ser Ala Ala Asn Lys Gly
145 150 155 160

Ser Leu Ser Cys Val Lys Glu His Thr Tyr Lys Val Glu Met Ser His 165 170 175

Asn Gln Ala Leu Gly Lys Val Asn Val Leu Ser Pro Asn Arg Asn Val 180 185 190

His Glu Trp Leu Tyr Ser Phe Lys Pro Asn Phe Asn Gln Ile Glu Ser 195 200 205

Asn Asn Arg Thr Val Asn Ser Leu Ala Val Lys Ser Leu Leu Met Ala 210 215 220

Thr Glu Asn Asn Ile Met Pro Asn Ser Gln Ala Phe Val Lys Ala Ser 225 230 235 240

Thr Asp Ser His Phe Lys Leu Ser Leu Trp Leu Arg Ile Pro Lys Val 245 250 255

Leu Lys Gln Ile Ala Ile Gln Lys Leu Phe Lys Phe Ala Gly Asp Glu

260 265 270

Thr Gly Lys Ser Phe Tyr Leu Ser Ile Ala Cys Ile Pro Asn His Asn 275 280 285

Ser Val Glu Thr Ala Leu Asn Val Thr Val Ile Cys Arg His Gln Leu 290 295 300

Pro Ile Pro Lys Ser Lys Ala Pro Phe Glu Leu Ser Met Ile Phe Ser 305 310 315 320

Asp Leu Lys Glu Pro Tyr Asn Thr Val His Asp Pro Ser Tyr Pro Gln 325 330 335

Arg Ile Val His Ala Leu Leu Glu Thr His Thr Ser Phe Ala Gln Val 340 345 350

Leu Cys Asn Lys Leu Gln Glu Asp Val Ile Ile Tyr Thr Ile Asn Ser 355 360 365

Pro Glu Leu Thr Pro Ala Lys Leu Asp Leu Gly Glu Arg Thr Leu Asn 370 375 380

Tyr Ser Glu Asp Ala Ser Lys Lys Lys Tyr Phe Leu Ser Lys Thr Leu 385 390 395 400

Glu Cys Leu Pro Val Asn Val Gln Thr Met Ser Tyr Leu Asp Ser Ile 405 410 415

Gln Ile Pro Ser Trp Lys Ile Asp Phe Ala Arg Gly Glu Ile Arg Ile 420 425 430

Ser Pro Gln Ser Thr Pro Ile Ala Arg Ser Leu Leu Lys Leu Asp Leu 435 440 445

Ser Lys Ile Lys Glu Lys Lys Ser Leu Thr Trp Glu Thr Ser Ser Tyr 450 455 460

Asp Leu Glu 465

(2) INFORMATION FOR SEQ ID NO:13:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 258 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Met Ser Lys Val Lys Leu Thr Lys Glu Asn Ile Val Ser Leu Leu Thr 1 5 10 15

Gln Ser Ala Asp Val Glu Phe Glu Glu Asp Gln Asn Gln Val Ala Phe 20 25 30

Asn Phe Lys Thr Phe Cys Gln Glu Asn Leu Asp Leu Ile Lys Lys Met 35 40 45

Ser Ile Thr Ser Cys Leu Thr Phe Leu Lys Asn Arg Gln Gly Ile Met 50 55 60

Lys Val Val Asn Gln Ser Asp Phe Thr Phe Gly Lys Val Thr Ile Lys 65 70 75 80

Lys Asn Ser Glu Arg Val Gly Ala Lys Asp Met Thr Phe Arg Arg Leu 85 90 95

Asp Ser Met Ile Arg Val Lys Leu Ile Glu Glu Thr Ala Asn Asn Glu 100 105 110

Asn Leu Ala Ile Ile Lys Ala Lys Ile Ala Ser His Pro Leu Val Gln
115 120 125

Ala Tyr Gly Leu Pro Leu Ala Asp Ala Lys Ser Val Arg Leu Ala Ile 130 135 140

Met Ile Ser Val Val Leu Ala Ile Tyr Gln Asp Ala Lys Tyr Lys Glu 165 170 175

Leu Gly Ile Glu Pro Thr Lys Tyr Asn Thr Lys Glu Ala Leu Gly Lys 180 185 190

Val Cys Thr Val Leu Lys Ser Lys Gly Phe Thr Met Asp Asp Ala Gln 195 200 205

Ile Asn Lys Gly Lys Glu Tyr Ala Lys Ile Leu Ser Ser Cys Asn Pro 210 215 220

Asn Ala Lys Gly Ser Ile Ala Met Asp Tyr Tyr Ser Asp Asn Leu Asp 225 230 235 240

Lys Phe Tyr Glu Met Phe Gly Val Lys Lys Glu Ala Lys Ile Ala Gly 245 250 255

Val Ala

(2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 3049 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

AGAGCAATTG	GGTCATTTTT	TATTCTAAAT	CGAACCTCAA	CTAGCAAATC	TCAGAACTGT	60
AATAAGCACA	AGAGCACAAG	AGCCACAATG	TCATCAGGTG	TTTATGAATC	GATCATTCAG	120
ACAAAGGCTT	CAGTTTGGGG	ATCGACAGCA	TCTGGTAAGT	CCATCGTGGA	TTCTTACTGG	180
ATTTATGAGT	TTCCAACTGG	TTCTCCACTG	GTTCAAACTC	AGTTGTACTC	TGATTCGAGG	240
AGCAAAAGTA	GCTTCGGCTA	CACTTCAAAA	ATTGGTGATA	TTCCTGCTGT	AGAGGAGGAA	300
ATTTTATCTC	AGAACGTTCA	TATCCCAGTG	TTTGATGATA	TTGATTTCAG	CATCAATATC	360
AATGATTCTT	TCTTGGCAAT	TTCTGTTTGT	TCCAACACAG	TTAACACCAA	TGGAGTGAAG	420
CATCAGGGTC	ATCTTAAAGT	TCTTTCTCTT	GCCCAATTGC	ATCCCTTTGA	ACCTGTGATG	480
AGCAGGTCAG	AGATTGCTAG	CAGATTCCGG	CTCCAAGAAG	AAGATATAAT	TCCTGATGAC	540
AAATATATAT	CTGCTGCTAA	CAAGGGATCT	CTCTCCTGTG	TCAAAGAACA	TACTTACAAA	600
GTCGAAATGA	GCCACAATCA	GGCTTTAGGC	AAAGTGAATG	TTCTTTCTCC	TAACAGAAAT	660
GTTCATGAGT	GGCTGTATAG	TTTCAAACCA	AATTTCAACC	AGATCGAAAG	TAATAACAGA	720
ACTGTAAATT	CTCTTGCAGT	CAAATCTTTG	CTCATGGCTA	CAGAAAACAA	CATTATGCCT	780
AACTCTCAAG	CTTTTGTTAA	AGCTTCTACT	GATTCTCATT	TTAAGTTGAG	CCTTTGGCTG	840
AGAATTCCAA	AAGTTTTGAA	GCAAATAGCC	ATACAGAAGC	TCTTCAAGTT	TGCAGGAGAC	900
GAAACCGGTA	AAAGTTTCTA	TTTGTCTATT	GCATGCATCC	CAAATCACAA	CAGTGTGGAA	960
ACAGCTTTAA	ATGTCACTGT	TATATGTAGA	CATCAGCTTC	CAATCCCTAA	GTCCAAAGCT	1020
CCTTTTGAAT	TATCAATGAT	TTTCTCCGAT	CTGAAAGAGC	CTTACAACAC	TGTGCATGAT	1080
CCTTCATATC	CTCAAAGGAT	TGTTCATGCT	TTGCTTGAGA	CTCACACTTC	CTTTGCACAA	1140
GTTCTCTGCA	ACAAGCTGCA	AGAAGATGTG	ATCATATATA	CTATAAACAG	CCCTGAACTA	1200
ACCCCAGCTA	AGCTGGATCT	AGGTGAAAGA	ACCTTGAACT	ACAGTGAAGA	TGCTTCGAAG	1260
AAGAAGTATT	TTCTTTCAAA	AACACTCGAA	TGCTTGCCAG	TAAATGTGCA	GACTATGTCT	1320
TATTTGGATA	GCATCCAGAT	TCCTTCATGG	AAGATAGACT	TTGCCAGAGG	AGAGATCAGA	1380

ATCTCCCCTC	AATCTACTCC	TATTGCAAGA	TCTTTGCTCA	AGCTGGATTT	GAGCAAGATC	1440
AAGGAAAAGA	AGTCCTTGAC	TTGGGAAACA	TCCAGCTATG	ATCTAGAATA	AAAGTGGCTC	1500
ATACTACTCT	AAGTAGTATT	TGTCAACTTG	CTTATCCTTT	ATGTTGTTTA	TTTCTTTTAA	1560
ATCTAAAGTA	AGTTAGATTC	AAGTAGTTTA	GTATGCTATA	GCATTATTAC	AAAAAATACA	1620
AAAAAATACA	AAAAAATACA	AAAAATATAA	AAAACCCAAA	AAGATCCCAA	AAGGGACGAT	1680
TTGGTTGATT	TACTCTGTTT	TAGGCTTATC	TAAGCTGCTT	TTGTTTGAGC	AAAATAACAT	1740
TGTAACATGC	AATAACTGGA	ATTTAAAGTC	CTAAAAGAAG	TTTCAAAGGA	CAGCTTAGCC	1800
AAAATTGGTT	TTTGTTTTTG	TTTTTTTTGTT	TTTTGTTTTT	TTGTTTTATT	TTTATTTTTA	1860
GTTTATTTTT	TGTTTTTGTT	ATTTTTATTT	TTATTTTATT	TTCTTTTATT	TTATTTATAT	1920
ATATATCAAA	CACAATCCAC	ACAAATAATT	TTAATTTCAA	ACATTCTACT	GATTTAACAC	1980
ACTTAGCCTG	ACTTTATCAC	ACTTAACACG	CTTAGTTAGG	CTTTAACACA	CTGAACTGAA	2040
TTAAAACACA	CTTAGTATTA	TGCATCTCTT	AATTAACACA	CTTTAATAAT	ATGCATCTCT	2100
GAATCAGCCT	TAAAGAAGCT	TTTATGCAAC	ACCAGCAATC	TTGGCCTCTT	TCTTAACTCC	2160
AAACATTTCA	TAGAATTTGT	CAAGATTATC	ACTGTAATAG	TCCATAGCAA	TGCTTCCCTT	2220
AGCATTGGGA	TTGCAAGAAC	TAAGTATCTT	GGCATATTCT	TTCCCTTTGT	TTATCTGTGC	2280
ATCATCCATT	GTAAATCCTT	TGCTTTTAAG	CACTGTGCAA	ACCTTCCCCA	GAGCTTCCTT	2340
AGTGTTGTAC	TTAGTTGGTT	CAATCCCTAA	CTCCTTGTAC	TTTGCATCTT	GATATATGGC	2400
AAGAACAACA	CTGATCATCT	CGAAGCTGTC	AACAGAAGCA	ATGAGAGGGA	TACTACCTCC	2460
AAGCATTATA	GCAAGTCTCA	CAGATTTTGC	ATCTGCCAGA	GGCAGCCCGT	AAGCTTGGAC	2520
CAAAGGGTGG	GAGGCAATTT	TTGCTTTGAT	AATAGCAAGA	TTCTCATTGT	TTGCAGTCTC	2580
TTCTATGAGC	TTCACTCTTA	TCATGCTATC	AAGCCTCCTG	AAAGTCATAT	CCTTAGCTCC	2640
AACTCTTTCA	GAATTTTTCT	TTATCGTGAC	CTTACCAAAA	GTAAAATCAC	TTTGGTTCAC	2700
AACTTTCATA	ATGCCTTGGC	GATTCTTCAA	GAAAGTCAAA	CATGAAGTGA	TACTCATTTT	2760
CTTAATCAGG	TCAAGATTTT	CCTGACAGAA	AGTCTTAAAG	TTGAATGCGA	CCTGGTTCTG	2820
GTCTTCTTCA	AACTCAACAT	CTGCAGATTG	AGTTAAAAGA	GAGACAATGT	TTTCTTTTGT	2880
GAGCTTGACC	TTAGACATGG	TGGCAGTTTA	GATCTAGACC	TTTCTCGAGA	GATAAGATTC	2940
AAGGTGAGAA	AGTGCAACAC	TGTAGACCGC	GGTCGTTACT	TATCCTGTTA	ATGTGATGAT	3000

(2) INFORMATION FOR SEQ ID NO:15:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 777 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

TTATGCAACA CCAGCAATCT TGGCCTCTTT CTTAACTCCA AACATTTCAT AGAATTTGTC 60 AAGATTATCA CTGTAATAGT CCATAGCAAT GCTTCCCTTA GCATTGGGAT TGCAAGAACT 120 AAGTATCTTG GCATATTCTT TCCCTTTGTT TATCTGTGCA TCATCCATTG TAAATCCTTT 180 GCTTTTAAGC ACTGTGCAAA CCTTCCCCAG AGCTTCCTTA GTGTTGTACT TAGTTGGTTC 240 AATCCCTAAC TCCTTGTACT TTGCATCTTG ATATATGGCA AGAACAACAC TGATCATCTC 300 GAAGCTGTCA ACAGAAGCAA TGAGAGGGAT ACTACCTCCA AGCATTATAG CAAGTCTCAC 360 AGATTTTGCA TCTGCCAGAG GCAGCCCGTA AGCTTGGACC AAAGGGTGGG AGGCAATTTT 420 TGCTTTGATA ATAGCAAGAT TCTCATTGTT TGCAGTCTCT TCTATGAGCT TCACTCTTAT 480 CATGCTATCA AGCCTCCTGA AAGTCATATC CTTAGCTCCA ACTCTTTCAG AATTTTTCTT 540 TATCGTGACC TTACCAAAAG TAAAATCACT TTGGTTCACA ACTTTCATAA TGCCTTGGCG 600 ATTCTTCAAG AAAGTCAAAC ATGAAGTGAT ACTCATTTTC TTAATCAGGT CAAGATTTTC 660 CTGACAGAAA GTCTTAAAGT TGAATGCGAC CTGGTTCTGG TCTTCTTCAA ACTCAACATC 720 TGCAGATTGA GTTAAAAGAG AGACAATGTT TTCTTTTGTG AGCTTGACCT TAGACAT 777

(2) INFORMATION FOR SEQ ID NO:16:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

18

660

(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID	NO:17:
TTATATCTTC TTCTTGGA	
(2) INFORMATION FOR SEQ ID NO:18:	
(2) INFORMATION FOR SEQ ID NO:18: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1401 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ I	5
(ii) MOLECULE TYPE: DNA (genomic)
(xi) SEQUENCE DESCRIPTION: SEQ I	
ATGTCATCAG GTGTTTATGA ATCGATCATT CAG	ACAAAGG CTTCAGTTTG GGGATCGACA 60
GCATCTGGTA AGTCCATCGT GGATTCTTAC TGG	
CTGGTTCAAA CTCAGTTGTA CTCTGATTCG AGC	GAGCAAAA GTAGCTTCGG CTACACTTCA 180
AAAATTGGTG ATATTCCTGC TGTAGAGGAG GA	AATTTTAT CTCAGAACGT TCATATCCCA 240
GTGTTTGATG ATATTGATTT CAGCATCAAT AT	CAATGATT CTTTCTTGGC AATTTCTGTT 300
TGTTCCAACA CAGTTAACAC CAATGGAGTG AA	GCATCAGG GTCATCTTAA AGTTCTTTCT 360
CTTGCCCAAT TGCATCCCTT TGAACCTGTG AT	GAGCAGGT CAGAGATTGC TAGCAGATTC 420
CGGCTCCAAG AAGAAGATAT AATTCCTGAT GA	
TCTCTCCT GTGTCAAAGA ACATACTTAC AA	
GGCAAAGTGA ATGTTCTTTC TCCTAACAGA AA	

CCAAATTTCA ACCAGATCGA AAGTAATAAC AGAACTGTAA ATTCTCTTGC AGTCAAATCT

GTTCTGAGAT TTGCTAGT

(2) INFORMATION FOR SEQ ID NO:17:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 18 base pairs (B) TYPE: nucleic acid

TTGCTCATGG	CTACAGAAAA	CAACATTATG	CCTAACTCTC	AAGCTTTTGT	TAAAGCTTCT	720
ACTGATTCTC	ATTTTAAGTT	GAGCCTTTGG	CTGAGAATTC	CAAAAGTTTT	GAAGCAAATA	780
GCCATACAGA	AGCTCTTCAA	GTTTGCAGGA	GACGAAACCG	GTAAAAGTTT	CTATTTGTCT	840
ATTGCATGCA	TCCCAAATCA	CAACAGTGTG	GAAACAGCTT	TAAATGTCAC	TGTTATATGT	900
AGACATCAGC	TTCCAATCCC	TAAGTCCAAA	GCTCCTTTTG	AATTATCAAT	GATTTTCTCC	960
GATCTGAAAG	AGCCTTACAA	CACTGTGCAT	GATCCTTCAT	ATCCTCAAAG	GATTGTTCAT	1020
GCTTTGCTTG	AGACTCACAC	TTCCTTTGCA	CAAGTTCTCT	GCAACAAGCT	GCAAGAAGAT	1080
GTGATCATAT	ATACTATAAA	CAGCCCTGAA	CTAACCCCAG	CTAAGCTGGA	TCTAGGTGAA	1140
AGAACCTTGA	ACTACAGTGA	AGATGCTTCG	AAGAAGAAGT	ATTTTCTTTC	AAAAACACTC	1200
GAATGCTTGC	CAGTAAATGT	GCAGACTATG	TCTTATTTGG	ATAGCATCCA	GATTCCTTCA	1260
TGGAAGATAG	ACTTTGCCAG	AGGAGAGATC	AGAATCTCCC	CTCAATCTAC	TCCTATTGCA	1320
AGATCTTTGC	TCAAGCTGGA	TTTGAGCAAG	ATCAAGGAAA	AGAAGTCCTT	GACTTGGGAA	1380
ACATCCAGCT	ATGATCTAGA	A				1401

(2) INFORMATION FOR SEQ ID NO:19:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 777 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

ATGTCTAAGG	TCAAGCTCAC	AAAAGAAAAC	ATTGTCTCTC	TTTTAACTCA	ATCTGCAGAT	60
GTTGAGTTTG	AAGAAGACCA	GAACCAGGTC	GCATTCAACT	TTAAGACTTT	CTGTCAGGAA	120
AATCTTGACC	TGATTAAGAA	AATGAGTATC	ACTTCATGTT	TGACTTTCTT	GAAGAATCGC	180
CAAGGCATTA	TGAAAGTTGT	GAACCAAAGT	GATTTTACTT	TTGGTAAGGT	CACGATAAAG	240
AAAAATTCTG	AAAGAGTTGG	AGCTAAGGAT	ATGACTTTCA	GGAGGCTTGA	TAGCATGATA	300
AGAGTGAAGC	TCATAGAAGA	GACTGCAAAC	AATGAGAATC	TTGCTATTAT	CAAAGCAAAA	360
ATTGCCTCCC	ACCCTTTGGT	CCAAGCTTAC	GGGCTGCCTC	TGGCAGATGC	AAAATCTGTG	420
AGACTTGCTA	TAATGCTTGG	AGGTAGTATC	CCTCTCATTG	CTTCTGTTGA	CAGCTTCGAG	480

ATGATCAGTG TTGTTCTTGC CATATATCAA GATGCAAAGT ACAAGGAGTT AGGGATTGAA	540
CCAACTAAGT ACAACACTAA GGAAGCTCTG GGGAAGGTTT GCACAGTGCT TAAAAGCAAA	600
GGATTTACAA TGGATGATGC ACAGATAAAC AAAGGGAAAG AATATGCCAA GATACTTAGT	660
TCTTGCAATC CCAATGCTAA GGGAAGCATT GCTATGGACT ATTACAGTGA TAATCTTGAC	720
AAATTCTATG AAATGTTTGG AGTTAAGAAA GAGGCCAAGA TTGCTGGTGT TGCATAA	777
(2) INFORMATION FOR SEQ ID NO:20: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 40 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
TACTTATCTA GAACCATGGA CAAAGCAAAG ATTACCAAGG	40
(2) INFORMATION FOR SEQ ID NO:21: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 42 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
TACAGTGGAT CCATGGTTAT TTCAAATAAT TTATAAAAGC AC	42
(2) INFORMATION FOR SEQ ID NO:22:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 36 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	

(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
AGCATTGGAT CCATGGTTAA CACACTAAGC AAGCAC	36
(2) INFORMATION FOR SEQ ID NO:23:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 46 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
AGCTAATCTA GAACCATGGA TGACTCACTA AGGAAAGCAT TGTTGC	46
(2) INFORMATION FOR SEQ ID NO:24:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
CCCACTATCC TTCGCAAGAC CC	22
(2) INFORMATION FOR SEQ ID NO:25:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
TACAGTGGAT CCATGGTTAA GGTAATCCAT AGGCTTGAC	39
(2) INFORMATION FOR SEQ ID NO:26:	

	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 40 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
AGC:	AACCAT GGTTAAGCTC ACTAAGGAAA GCATTGTTGC	40
(2)	INFORMATION FOR SEQ ID NO:27:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 46 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
AGC'	AATCTA GAACCATGGA TGACTCACTA AGGAAAGCAT TGTTGC	46
(2)	INFORMATION FOR SEQ ID NO:28:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 36 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
AGC	TTGGAT CCATGGTTAA CACACTAAGC AAGCAC	36
(2)	INFORMATION FOR SEQ ID NO:29:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECILE TYPE: DNA (genomic)	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
TACAGTTCTA GAACCATGGA TGATGCAAAG TCTGTGAGG	39
(2) INFORMATION FOR SEQ ID NO:30:	
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 49 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
AGATTCTCTA GACCATGGTG ACTTGATGAG CAAAGTCTGT GAGGCTTGC	49